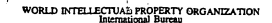
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(54) Title: NEW STAPHYLOKINASE DERIVATIVES

(57) Abstract

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A method for producing the derivatives of the invention by preparing a DNA fragment comprising at least the part of the coding sequence of staphylokinase that provides for its biological activity; performing in vitro site-directed mutagenesis on the DNA fragment to replace one or more codons for wild-type amino acids by a codon for another amino acid; cloning the mutated DNA fragment in a suitable vector, transforming or transfecting a suitable host cell with the vector, and culturing the host cell under conditions suitable for expressing the DNA fragment. Preferably the DNA fragment is a 453 bp EcoRI-HindIII fragment of the plasmid pMEX602SakB, the in vitro site-directed mutagenesis is performed by an oligonucleotide-directed mutagenesis system using the plasmid pMa/c and the repair deficient E. coli strain WK6MutS, and the mutated DNA fragment is expressed E. coli strain WK6. The invention also relates to pharmaceutical compositions comprising at least one of the staphylokinase derivatives according to the invention together with a suitable excipient, for treatment of arterial thrombosis.

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(74) Agent: VAN SOMEREN, Petronella, Francisca, Hendrika, Maria; Arnold & Siedsma, Sweelinckplein 1, NL-2517 GK The Hague (NL).

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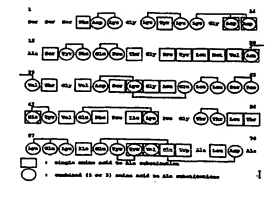
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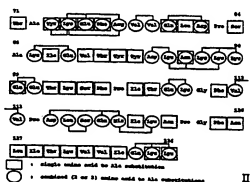
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(54) Title: IDENTIFICATION, PRODUCTION AND USE OF STAPHYLOKINASE DERIVATIVES WITH REDUCED IMMUNO-GENICITY AND/OR REDUCED CLEARANCE

(57) Abstract

Methods for the identification, production and use of staphylokinase derivatives characterized by a reduced immunogenicity after administration in patients and that can be administered by single intravenous bolus injection. The derivatives of the invention are obtained by preparing a DNA fragment comprising at least the part of the coding sequence of staphylokinase that provides for its biological activity; performing in vitro site-directed mutagenesis on the DNA fragment to replace one or more codons for wild-type amino acids by a codon for another amino acid; cloning the mutated DNA fragment in a suitable vector, transforming or transfecting a suitable host cell with the vector, culturing the host cell under conditions suitable for expressing the DNA fragment; purifying the expressed staphylokinase derivative to homogeneity and chemically modifying substituted Cys residues with thiol-directed polyethylene glycol; preferably the DNA fragment is a 453 bp EcoRI-HindIII fragment of the plasmid pMEX602sakB, (pMEX.SakSTAR), the in vitro site-directed mutagenesis is performed by spliced overlap extension polymerase chain reaction and the mutated DNA fragment is expressed in E. coli strain TG1 or WK6. The invention also relates to pharmaceutical compositions comprising at least one of the staphylokinase derivatives according to the invention together with a suitable excipient, for treatment of arterial thrombosis.





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WO 99/40198 PCT/EP99/00748

IDENTIFICATION, PRODUCTION AND USE OF STAPHYLOKINASE DERIVATIVES WITH REDUCED IMMUNOGENICITY AND/OR REDUCED CLEARANCE

5 The present invention relates to new staphylokinase derivatives with reduced immunogenicity which can be administered by continuous infusion or by single intravenous bolus injection, to their identification, production and use in the treatment of arterial thrombosis and to the preparation of a pharmaceutical composition for treating arterial thrombosis. More in particular the invention relates to the use of engineered staphylokinase derivatives for the preparation of a pharmaceutical composition for treating myocardial infarction.

Staphylokinase, a protein produced by certain strains of Staphylococcus aureus, which was shown to have profibrinolytic properties more than 4 decades ago (1, 2) appears to constitute a potent thrombolytic agent in 20 patients with acute myocardial infarction (3, 4). The staphylokinase gene has been cloned from the bacteriophages sak\(operation C \) and sak42D (6) as well as from the genomic DNA (sakSTAR) of a lysogenic Staphylococcus aureus strain (7). The staphylokinase gene encodes a 25 protein of 163 amino acids, with amino acid 28 corresponding to the NH,-terminal residue of full length mature staphylokinase (6, 8, 9). The mature protein sequence of the wild-type variant SakSTAR (9) is represented in Figure 1. Only four nucleotide differences 30 were found in the coding regions of the sak ϕ C, sak42D and sakSTAR genes, one of which constituted a silent mutation (6, 8, 9). In a plasma milieu, staphylokinase is able to dissolve fibrin clots without associated fibrinogen degradation (10-12). This fibrin-specificity of 35 staphylokinase is the result of reduced inhibition by α_2 -antiplasmin of plasmin.staphylokinase complex bound to fibrin, recycling of staphylokinase from the plasmin.staphylokinase complex following inhibition by

 α_2 -antiplasmin, and prevention of the conversion of circulating plasminogen.staphylokinase to plasmin.staphylokinase by α_2 -antiplasmin (13-15). In addition staphylokinase has a weak affinity for circulating but a

- 5 high affinity for fibrin-bound plasminogen (16) and staphylokinase requires NH₂-terminal processing by plasmin to display its plasminogen activating potential (17). In several experimental animal models, staphylokinase appears to be equipotent to streptokinase for the
- 10 dissolution of whole blood or plasma clots, but significantly more potent for the dissolution of platelet-rich or retracted thrombi (18, 19).

 Staphylokinase is a heterologous protein and is immunogenic in man. The intrinsic immunogenicity of
- 15 staphylokinase, like that of streptokinase, clearly hampers its unrestricted use. Not only will patients with preexisting high antibody titers be refractory to the thrombolytic effect of these agents, but allergic side effects and occasional life-threatening anaphylaxis may
- 20 occur (20). Because both streptokinase and staphylokinase are heterologous proteins, it is not obvious that their immunogenicity could be reduced by protein engineering. Indeed, no successful attempts to generate active low molecular weight fragments from streptokinase have been
- 25 reported. In staphylokinase, deletion of the NH₂-terminal 17 amino acids or the COOH-terminal 2 amino acids inactivates the molecule, which in addition is very sensitive to inactivation by site-specific mutagenesis (21).
- It is therefore the object of the present invention to provide less immunogenic variants of staphylokinase having preferably a higher specific activity and/or lower plasma clearance and/or increased thrombolytic potency.
- In the research that ultimately led to the present invention it was already found that the wild-type staphylokinase variant SakSTAR (9) contains three non-overlapping immunodominant epitopes, at least two of

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which can be eliminated by specific site-directed mutagenesis, without inactivation of the molecule. This has been disclosed in EP-95200023.0 (22). These engineered staphylokinase variants are less reactive with antibodies elicited in patients treated with wild-type staphylokinase, and are significantly less immunogenic than wild-type staphylokinase, as demonstrated in rabbit and baboon models and in patients with peripheral arterial occlusion (22).

10 The present invention now relates to general methods for the identification, production and use of staphylokinase derivatives showing a reduced antigenicity and immunogenicity as compared to wild-type staphylokinase as well as for variants with selective 15 derivatization with polyethylene glycol. The derivatives preferably have a higher specific activity and/or lower plasma clearance and/or increased thrombolytic potency. The derivatives have essentially the amino acid sequence of wild-type staphylokinase or modified versions thereof 20 and essentially intact biological activities, but have a reduced reactivity with a panel of murine monoclonal antibodies and/or with antibodies induced in patients by treatment with wild-type SakSTAR. The polyethylene glycol substituted ("pegylated") variants have reduced plasma 25 clearances rendering them particularly suited for use by single intravenous bolus administration. Instead of PEG other pharmaceutically acceptable macromolecules can be used.

More in particular, the invention provides for staphylokinase derivatives SakSTAR(K35X,G36X,E65X,K74X,E80X,D82X,K102X,E108X,K109X,K121X,K130X,K135X,K136X,+137X) having the amino acid sequence as depicted in figure 1 in which the amino acids Lys in position 35, Gly in position 36, Glu in position 65, Lys in position 74, 35 Glu in position 80, Asp in position 82, Lys in position 102, Glu in position 108, Lys in position 109, Lys in position 121, Lys in position 130, Lys in position 135 and/or Lys in position 136 have been replaced with other amino acids and/or in which one amino acid has been added 40 at the COOH-terminus, thus altering the immunogenicity

after administration in patients, without markedly reducing the specific activity.

Further preferred embodiments of the invention are staphylokinase derivatives listed in Tables 1, 3, 4, 5, 6, 7, 8, 13, 19 and 20, having the amino acid sequence as depicted in figure 1 in which the indicated amino acids have been replaced by other amino acids thus reducing the absorption of SakSTAR-specific antibodies from plasma of patients treated with staphylokinase, without reducing the specific activity.

Derivatives in which the specific activity is increased and the immunogenicity is decreased are the following:

SakSTAR(K74A, E75A, R77A), SakSTAR(K35A, E75A),

- 15 SakSTAR(E75A), SakSTAR(E80A,D82A), SakSTAR(E80A),
 SakSTAR(D82A), SakSTAR(E75A,D82A), SakSTAR(S34G,G36R,
 H43R), SakSTAR(K35A), SakSTAR(E80A), SakSTAR(D82A,S84A),
 SakSTAR(T90A), SakSTAR(Y92A), SakSTAR(K130A),
 SakSTAR(V132A), SakSTAR(S34G,G36R,H43R), SakSTAR(G36R),
- 20 SakSTAR(H43R), SakSTAR(G36R,K74R), SakSTAR(K35E),
 SakSTAR(K74Q), SakSTAR(K130T), SakSTAR(V132L),
 SakSTAR(V132T), SakSTAR(V132N), SakSTAR(V132R),
 SakSTAR(K130T,K135R), SakSTAR(G36R,K130T,K135R),
 SakSTAR(K74R,K130T,K135R), SakSTAR(K74Q,K130T,K135R),
- 25 SakSTAR(G36R, K74R, K130T, K135R), SakSTAR(G36R, K74Q, K130T, K135R), SakSTAR(G36R, H43R, K74R, K130T, K135R), SakSTAR(E65A, K74Q, K130T, K135R), SakSTAR(E65Q, K74Q, K130T, K135R), SakSTAR(K74Q, K86A, K130T, K135R), SakSTAR(E65Q, T71S, K74Q, K130T, K135R),
- 30 SakSTAR(K74Q,K130A,K135R), SakSTAR(E65Q,K74Q,K130A, K135R), SakSTAR(K74Q,K130E,K135R), SakSTAR(K74Q,K130E, V132R,K135R), SakSTAR(E65Q,K74Q,T90A,K130A,K135R), SakSTAR(E65Q,K74Q,N95A,K130A,K135R), SakSTAR(E65Q,K74Q, E118A,K130A,K135R), SakSTAR(E65Q,K74Q,N95A,E118A,K130A,
- 35 K135R), SakSTAR(N95A,K130A,K135R), SakSTAR(E65Q,K74Q, K109A,K130,K135R), SakSTAR(E65Q,K74Q,E108A,K109A,

K130T,K135R), SakSTAR(E65Q,K74Q,K121A,K130T,K135R),
SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R,K136A,+137K),
SakSTAR(E80A,D82A,K130T,K135R), SakSTAR(K74R,E80A,D82A,
K130T,K135R), SakSTAR(K74Q,E80A,D82A,K130T,K135R),
SakSTAR(K36A,K74R,E80A,D82A,K130T,K135R),
SakSTAR(K36A,K74R,E80A,D82A,K130T,K135R), SakSTAR(F65R)

- 5 SakSTAR(K35A, K74R, E80A, D82A, K130T, K135R), SakSTAR(E65D, K74R, E80A, D82A, K130T, K135R), SakSTAR(E65S, K74R, E80A, D82A, K130T, K135R), SakSTAR(S34G, G36R, K74R, K130T, K135R), SakSTAR(E65A, K74R, E80A, D82A, K130T, K135R), SakSTAR(E65N, K74R, E80A, D82A, K130T, K135R), SakSTAR(E65Q, K74R, E80A,
- 10 D82A,K130T,K135R), SakSTAR(K57A,E58A,E61A,E80A,D82A,
 K130T,K135R), SakSTAR(E65D,K74Q,E80A,D82A,K130T,K135R),
 SakSTAR(E65Q,K74Q,E80A,D82A,K130T,K135R), SakSTAR(K35A,
 E65D,K74Q,E80A,D82A,K130T,K135R), SakSTAR(K74R,E80A,D82A,
 S103A,K130T,K135R), SakSTAR(E65D,K74R,E80A,D82A,K109A,

(::::/

15 K130T,K135R), SakSTAR(E65D,K74R,E80A,D82A,K130T, K135R,K136A), SakSTAR(E65Q,K74Q,D82A,S84A,K130T,K135R), SakSTAR(K35A,K74Q,E80A,D82A,K130T,K135R), SakSTAR(K35A, E65D,K74R,E80A,D82A,K130T,K135R).

Of these SakSTAR(E65D,K74R,E80A,D82A,K130T,
20 K135R) having the code SY19 and SakSTAR(K35A,E65Q,K74R,
E80A,D82A,T90A,E99D,T101S,E108A,K109A,K130T,K135R) having
the code SY161 are especially preferred.

Besides the above described substitution derivatives the invention relates to derivatives having in addition an amino acid substituted with Cys. This type of substitution may result in dimerization and/or increased specific activity and/or reduced clearance and/or increased thrombolytic potency. Reduced plasma clearance is in particular obtained when the derivative is substituted with polyethylene glycol.

Preferred embodiments of such staphylokinase derivatives are those wherein the Cys is chemically modified with polyethylene glycol with molecular weights up to 20 kDa. In particular embodiments selected amino acids in the NH2-terminal region of 10 amino acids, are substituted with Cys, which is chemically modified with polyethylene glycol with molecular weights up to 20 kDa. These derivatives are characterized by a significantly

(A)

(E)

reduced plasma clearance and maintained thrombolytic potency upon single intravenous bolus administration at a reduced dose.

More in particular the serine in position 2 or 5 3 is substituted with a cystein and the cystein is chemically modified with polyethylene glycol having a molecular weight of 5, 10 or 20 kDa. Preferred embodiments of these derivatives are SY161(S3C-MP5), SY161(S3C-P10), SY161(S3C-P20), SY19(S3C-MP5), SY19(S3C-P10) all as defined in table 20.

The presence of cysteins allows the formation of dimers of two staphylokinase derivatives of the invention.

The invention also relates to a method for producing the derivatives of the invention by preparing a DNA fragment comprising at least the part of the coding sequence of staphylokinase that provides for its biological activity; performing in vitro site-directed 20 mutagenesis on the DNA fragment to replace one or more codons for wild-type amino acids by a codon for another amino acid; cloning the mutated DNA fragment in a suitable vector; transforming or transfecting a suitable host cell with the vector; culturing the host cell under 25 conditions suitable for expressing the DNA fragment; purifying the expressed staphylokinase derivative to homogeneity and derivatizing the variant with polyethylene glycol.

Preferably the DNA fragment is a 453 bp

30 EcoRI-HindIII fragment of the plasmid pMEX602sakB (22, 23), the <u>in vitro</u> site-directed mutagenesis is preferably performed by spliced overlap extension polymerase chain reaction. Such overlap extension PCR is preferably performed with Vent DNA polymerase (New England Biolabs)

35 or Taq polymerase (Boehringer Mannheim) and with available or generated wildtype sakSTAR or sakSTAR variants as template (24).

The invention also relates to pharmaceutical compositions comprising at least one of the staphylokinase derivatives according to the invention together with a suitable excipient, for treatment of 5 arterial thrombosis. Pharmaceutical compositions, containing less immunogenic staphylokinase variants or "pegylated" staphylokinase variants as the active ingredient, for treating arterial thrombosis in human or veterinary practice may take the form of powders or 10 solutions and may be used for intravenous, intraarterial or parenteral administration. Such compositions may be prepared by combining (e.g. mixing, dissolving etc.) the active compound with pharmaceutically acceptable excipients of neutral character (such as aqueous or 15 non-aqueous solvents, stabilizers, emulsifiers, detergents, additives), and further, if necessary with dyes.

Furthermore the invention relates to the use of the staphylokinase derivatives for the treatment of arterial thrombosis, in particular myocardial infarction, and to the use of staphylokinase derivatives for the preparation of a pharmaceutical composition for the treatment of arterial thrombosis, in particular myocardial infarction. In the above and the following the terms "derivatives", "mutants" and "variants" are used interchangeably.

Based on the present invention other variants and improvements will be obvious for the person skilled in the art. Thus random mutagenesis is likely to generate alternative mutants with reduced immunogenicity and possibly increased functional activity, whereas deletions or substitution with other amino acids may yield additional variants with reduced immunogenicity.

The present invention will be demonstrated in 35 more detail in the following examples, that are however not intended to be limiting to the scope of the invention. In the Examples reference is made to the following figures:

Fig 1. Protein sequence of wild-type staphylokinase, SakSTAR. Numbering starts with the NH2-terminal amino acid of mature full length staphylokinase.

Fig 2. Time course of neutralizing activities (left panel) and specific IgG against administered agent (right panel) following intra-arterial infusion of SakSTAR (open circles, n= 9), SakSTAR(K74A) (closed circles, n= 11) or SakSTAR(K74A,E75A, R77A) (open squares, n= 6) in patients with peripheral arterial occlusion. The data represent median values and

Fig 3. Protein sequence of wild-type staphylokinase, SakSTAR with indicated amino acid 15 substitutions.

interquartile ranges, in µg/mL.

squares: single amino acid substitutions; circles: combined (2 to 3) amino acid to Ala substitutions.

Fig. 4. Temperature stability of SakSTAR, (A);

20 SakSTAR(K74Q,E80A,D82A,K130T, K135R) (B);
SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R), (C); and
SakSTAR(K35A,E65D,K74Q,E80A,D82A, K130T,K135R), (D).
(○): 4°C; (●): 20°C; (▽): 37°C; (▼): 56°C; (□): 70°C.

Fig 5. Time course of neutralizing activities
25 (left panel) and specific IgG against administered agent
(right panel) following intra-arterial infusion of
SakSTAR (circles, n= 6),

SakSTAR(K74Q,E80A,D82A,K130T,K135R) (squares, n=6) or SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R) (triangles, n=6)

30 6) in patients with peripheral arterial occlusion. The data represent median values and 15-85 percentile ranges, in $\mu g/mL$.

EXAMPLES

35 EXAMPLE 1

Epitope mapping of wild-type staphylokinase

The epitope specificity of a panel of 15 murine MAbs (22) raised against wild-type SakSTAR was determined

(3)

by real-time biospecific interaction analysis (BIA) with the BIAcore instrument (Pharmacia, Biosensor AB, Uppsala, Sweden). The MAbs were immobilized on the surface of the Sencor Chip CM5 with the Amine Coupling Kit (Pharmacia 5 Biosensor AB) as recommended by the manufacturer (25). Immobilization was performed from protein solutions at a concentration of 20 μ g/mL in 10 mmol/L sodium acetate at pH 5.0 at a flow rate of 5 μ L/min during 6 minutes. This resulted in covalent attachment of 5,000 to 10,000 10 resonance unit (RU) of antibody (corresponding to 0.035 to 0.07 pmol/mm²). The SakSTAR solutions were passed by continuous flow at 20°C past the sensor surface. At least four concentrations of each analyte (range, 50 nmol/L to 50 mol/L) in 10 mmol/L HEPES, 3.4 mmol/L EDTA, 0.15 mol/L 15 NaCl, and 0.005% Surfactant P20, pH 7.2, were injected at a flow rate of 5 μ L/min during 6 minutes in the association phase. Then sample was replaced by buffer, also at a flow rate of 5 μ L/min during 6 minutes. After each cycle, the surface of the sensor chip was 20 regenerated by injection of 5 μ L of 15 mmol/L HCl. Apparent association (kass) and apparent dissociation

Apparent association (k_{ess}) and apparent dissociation (k_{diss}) rate constants were derived from the sensorgrams as described in detail elsewhere (26), and association equilibrium constants (K_A) calculated as their ratio.

Determination of the equilibrium association

constants for the binding of wild-type and variant
SakSTAR to insolubilized MAbs (Table 1) yielded apparent
association constants of 10⁷ to 10⁸ (mol/L)⁻¹, which are
one to two orders of magnitude lower than the apparent
30 association constants previously obtained for the binding
of these MAbs to insolubilized wild-type SakSTAR (22). If
the MAbs instead of the SakSTAR variants are
insolubilized, avidity effects of the bivalent MAbs are
avoided. The present values are indeed in better
35 agreement with known association constants of Mabs, and
therefore this "reversed" procedure was used throughout
the present invention.

In the tables the column indicated with "Variant" states the various staphylokinase derivatives which are identified by listing between brackets the substituted amino acids in single letter symbols followed 5 by their position number in the mature staphylokinase sequence and by the substituting amino acids in single letter symbol; the column "Exp." indicates expression levels in mg/L, and the column "Spec. Act." indicates the specific activity in Home Units as defined in example 2. 10 Indications "17G11", "26A2" etc. refer to monoclonal antibodies binding to the indicated epitopes I, II and III as defined in reference 22. Epitope I is recognized by the antibody cluster 17G11, 26A2, 30A2, 2B12 and 3G10, whereas epitope II is recognized by the antibody cluster 15 18F12, 14H5, 28H4, 32B2 and 7F10, and epitope III by the antibody cluster 7H11, 25E1, 40C8, 24C4 and 1A10. Human plasma "Pool" indicates a plasma pool from initially 16 and subsequently 10 patients immunized by treatment with

- SakSTAR, "Subpool B" indicates a plasma pool from three 20 patients that absorbed less than 50% of the induced antibodies with SakSTAR(K35A,E38A,K74A,E75A,R77A) and "Subpool C" indicates a plasma pool from 3 patients that absorbed >90% of the induced antibodies with SakSTAR(K35A,E38A,K74A,E75A,R77A) (22).
- In tables 6, 7 and 8 an additional pool of plasma from 40 patients immunized by treatment with SakSTAR (Pool 40) was also used.

EXAMPLE 2

30 Construction, epitope mapping with murine monoclonal antibodies and absorption with pooled plasma of immunized patients, of "alanine-to-wild-type" reversal variants of "charged-cluster-to-alanine" mutants of staphylokinase

1. <u>Introduction</u>

As stated above, wild-type staphylokinase
(SakSTAR variant (9)) contains three non-overlapping
immunodominant epitopes, two of which can be eliminated
by specific site-directed substitution of clusters of two

(K35A,E38A or E80A,D82A) or three (K74A,E75A,R77A) charged amino acids with Ala (22). The combination mutants SakSTAR(K35A,E38A,K74A,E75A,R77A) in which Lys35, Glu38, Lys74, Glu75 and Arg77, and SakSTAR(K74A,E75A,

- 5 R77A,E80A,D82A) in which Lys74, Glu75, Arg77, Glu80 and Asp82 were substituted with Ala (previously identified as SakSTAR.M3.8 and SakSTAR.M8.9, respectively (22)), were found to have a reduced reactivity with murine monoclonal antibodies against two of the three immunodominant
- 10 epitopes and to absorb on average only 2/3 of the neutralizing antibodies elicited in 16 patients by treatment with wild-type SakSTAR (22). These mutants also induced less antibody formation than wild-type SakSTAR in experimental thrombolysis models in rabbits and baboons,
- 15 and in patients with peripheral arterial occlusion (22).

 However, their specific activities were reduced to
 approximately 50% of that of wild-type SakSTAR, which
 would be of some concern with respect to the clinical use
 of these compounds.
- In an effort to improve the activity and stability without loss of the reduced antibody recognition, the effect of a systematic reversal of one or more of these substituted amino acids to the wild-type residues was studied. Fourteen new mutants were
- 25 constructed, purified and characterized in terms of specific activity, reactivity with the panel of murine monoclonal antibodies, and absorption of antibodies from plasma of patients treated with wild-type SakSTAR (Table 1). The present example thus focusses on reversal from
- 30 alanine to the wild-type residue of one or more of the seven amino acids of SakSTAR listed above i.e. K35, E38, K74, E75, R77, E80 and D82.

2. Reagents and Methods

35 The source of all reagents used in the present study has previously been reported (22). Restriction enzymes were purchased from Pharmacia (Uppsala, Sweden) or Boehringer Mannheim (Mannheim, Germany). T4 DNA

ligase, Klenow Fragment of <u>E. coli</u> DNA polymerase I and alkaline phosphatase were obtained from Boehringer Mannheim. Enzyme reactions were performed using the conditions suggested by the suppliers. Plasmid DNA was

- 5 isolated using a QIAGEN-purification protocol (provided by Westburg, Leusden, The Netherlands). pMEX.602sakB (i.e. pMEX.SakSTAR) was constructed as described elsewhere (23). SakSTAR, SakSTAR(K35A,E38A), SakSTAR(K74A,E75A,R77A), SakSTAR(E80A,D82A),
- 10 SakSTAR(K35A,E38A,K74A,E75A,R77A) and SakSTAR(K74A,E75A,R77A,E80A,D82A) were produced and purified
 as described elsewhere (22). Transformations of E. coli
 were performed utilizing the calcium phosphate procedure.
 DNA sequencing was performed using the dideoxy chain
- 15 termination reaction method and the Automated Laser fluorescent A.L.F. TM (Pharmacia). The chromogenic substrate (S2403) L-Pyroglutamyl-L-phenylalanyl-L-lysine-p-nitroanaline hydrochloride was purchased from Chromogenix (Belgium). 125I-labeled fibrinogen was
- 20 purchased from Amersham (UK). All other methods used in the present example have been previously described (22,27).

3. Construction of expression plasmids

- The plasmids encoding SakSTAR(K35A,E38A,K74A, E75A), SakSTAR(E38A,E75A,R77A), SakSTAR(E38A,E75A), SakSTAR(K35A,E75A,R77A), SakSTAR(K35A,E75A), SakSTAR(E80A), SakSTAR(D82A), SakSTAR(E75A,D82A), SakSTAR(K74A) and SakSTAR(E75A) were constructed by the
- 30 spliced overlap extension polymerase chain reaction (SOE-PCR) (24), using Vent DNA polymerase (New England Biolabs, Leusden, The Netherlands), and available or generated sakSTAR variants as template. Two fragments were amplified by PCR, the first one starting from the 5'
- 35 end of the staphylokinase gene with primer
 5'-CAGGAAACAGAATTCAGGAG-3' to the region to be
 mutagenized (forward primer), the second one from the
 same region (backward primer) to the 3' end of the

staphylokinase gene with primer 5'-CAAAACAGCCAAGCTTCATTCATTCAGC-3'. The forward and backward primers shared an overlap of around 24 bp (primers not shown). The two purified fragments were then 5 assembled together in a new primerless PCR using Tag polymerase (Boehringer Mannheim). After 7 cycles (1 min at 94°C, 1 min at 70°C), the extended product was reamplified by adding the 5' and 3' end primers (see above) to the PCR reaction and by cycling 25 times (1 min 10 at 94°C, 1 min 55°C, 1 min at 72°C). The final product was purified, digested with EcoRI and HindIII and cloned into the corresponding sites of pMEX602sakB. The plasmid encoding SakSTAR(E38A, K74A, E75A, R77A) was assembled by digestion of pMEX602sakB and pMEX.SakSTAR(K35A,E38A, 15 K74A,E75A,R77A) with BpmI which cuts between the codons for K35 and E38 of SakSTAR, and ligation of the required fragments. The plasmid encoding SakSTAR(K35A,K74A,E75A, R77A) was constructed by digestion of pMEX.SakSTAR(K35A, E38A, K74A, E75A, R77A) and pMEX.SakSTAR(K74A, E75A, R77A) 20 with BpmI and religation of the required fragments. The plasmids encoding SakSTAR(K35A, E38A, E75A, R77A) and SakSTAR(K35A,E38A,K74A,R77A) were constructed by two PCR using pMEX.SakSTAR(K35A,E38A,K74A,E75A,R77A) as template, followed by restriction ligation and recloning into 25 pMEX602sakB.

The SakSTAR variants were expressed and purified, as described below, from transformed E. coli

WK6 grown either in LB medium [SakSTAR(E38A,K74A,E75A, R77A), SakSTAR(K74A), SakSTAR(E75A) and SakSTAR(E75A, D82A)], or in terrific broth (TB) (28) medium [SakSTAR(K35A,K74A,E75A,R77A), SakSTAR(K35A,K74A,E75A,R77A), SakSTAR(K35A,E38A,E75A,R77A), SakSTAR(K35A,E38A,E75A,R77A), SakSTAR(K35A,E38A,E75A), SakSTAR(K35A,E75A), SakSTAR(K35A,E75A), SakSTAR(K35A,E75A), SakSTAR(K35A,E75A), SakSTAR(K35A,E75A), SakSTAR(K35A,E75A), SakSTAR(K35A,E75A),

SakSTAR(E80A), and SakSTAR(D82A)].

aliquot of an overnight saturated culture was used to inoculate a 2 L volume of LB medium containing 100 g/mL ampicillin. After 3 hours incubation at 37°C, IPTG (200 5 mol/L) was added to induce expression from the tac promoter. The production phase was allowed to proceed for 4 hours, after which the cells were pelleted by centrifugation at 4,000 rpm for 20 min, resuspended in 1/20 volume (100 mL) of 0.01 mol/L phosphate buffer pH 10 6.5 and disrupted by sonication at 0°C. Cell debris were removed by centrifugation for 20 min at 20,000 rpm and the supernatant, containing the cytosolic soluble protein fraction, was stored at -20°C until purification.

For the derivatives produced in TB medium, a 4 15 mL aliquot of an overnight saturated culture in LB medium was used to inoculate a 2 L culture in terrific broth containing 100 μ g/mL ampicillin. The culture was grown with vigorous aeration for 20 hours at 30°C. The cells were pelleted by centrifugation, resuspended in 1/10 20 volume (200 mL) of 0.01 mol/L phosphate buffer pH 6.5 and disrupted by sonication at 0°C. The suspension was then centrifuged for 20 min at 20,000 rpm and the supernatant was stored at -20°C until purification. Cleared cell lysates containing the SakSTAR variants were subjected to 25 chromatography on a 1.6 x 6 cm column of SP-Sephadex, followed by chromatography on a 1.6 x 5 cm column of Q-Sepharose [variants SakSTAR(E38A, K74A, E75A, R77A), SakSTAR(K35A, K74A, E75A, R77A), SakSTAR(K35A, E38A, E75A, R77A), SakSTAR(K35A, E38A, K74A, R77A) and

30 SakSTAR(K35A, E38A,K74A,E75A)] or by chromatography on a 1.6 x 6 cm column of phenyl-Sepharose [variants Sak-STAR(E35A,E38A,R77A), SakSTAR(E38A,E75A), SakSTAR-(K35A,E75A,R77A), SakSTAR(K35A,E75A), SakSTAR(K74A), SakSTAR(E75A), SakSTAR(E80A), SakSTAR(D82A) and Sak-

35 STAR(E75A,D82A)]. The SakSTAR containing fractions, localized by SDS-gel electrophoresis, were pooled for further analysis.

5. Physicochemical and biochemical analysis

Protein concentrations were determined according to Bradford (29). The specific activities of SakSTAR solutions were determined with a chromogenic 5 substrate assay carried out in microtiter plates using a mixture of 80 μ L SakSTAR solution and 100 μ L Glu-plasminogen solution prepared as described elsewhere (30) (final concentration 0.5 \(\mu\text{mol/L}\). After incubation for 30 min at 37°C, generated plasmin was quantitated by 10 addition of 20 µL S2403 (final concentration 1 mmol/L) and measurement of the absorption at 405 nm. The activity was expressed in home units (HU) by comparison with an in-house standard (lot STAN5) which was assigned an activity of 100,000 HU (100 kHU) per mg protein as 15 determined by amino acid composition (7). SDS-PAGE was performed with the Phast System (Pharmacia, Uppsala, Sweden) using 10-15% gradient gels and Coomassie Brilliant blue staining. Reduction of the samples was performed by heating at 100°C for 3 min in the presence 20 of 1% SDS and 1% dithiothreitol. The specific activities of the different SakSTAR mutants determined with the chromogenic substrate assay are summarized in Table 1.

6. Binding to murine monoclonal antibodies

25 In agreement with previous observations (22), SakSTAR(K74A,E75A,R77A) did not react with 4 of the 5 MAbs recognizing epitope I, whereas SakSTAR(K35A,E38A) did not react with 3 of the 5 and SakSTAR(E80A, D82A) not with 4 of the 5 Mabs recognizing epitope III. These 30 reduced reactivities were additive in SakSTAR(K35A,E38A, K74A, E75A, R77A) and in SakSTAR(K74A, E75A, R77A, E80A, D82A). The reduced reactivity of SakSTAR(K74A,E75A, R77A) was fully maintained in SakSTAR(K35A,E38A,K74A,E75A) and in SakSTAR(K35A, E75A,R77A), largely in SakSTAR(K35A,E38A, 35 E75A,R77A), SakSTAR(E38A,E75A,R77A), SakSTAR(E38A,E75A) and SakSTAR(E75A), but much less in SakSTAR(K35A,E38A, K74A,R77A) and SakSTAR(K74A), indicating that E75 is the

main contributor to the binding of the 4 Mabs recognizing

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epitope I of SakSTAR. However, surprisingly, binding of epitope I antibodies to SakSTAR(E75A,D82A) was normal in two independent preparations from expression plasmids with confirmed DNA sequences. The reduced reactivity of 5 the 3 MAbs of epitope III with SakSTAR(K35A,E38A) required both K35 and E38, as demonstrated with SakSTAR(E38A,K74A,E75A,R77A) and SakSTAR(K35A,K74A,E75A,R77A), with SakSTAR(E38A,E75A) and SakSTAR(K35A,E75A) and with SakSTAR(E38A,E75A,R77A) and SakSTAR(K35A,E75A,R77A).

10 The reduced reactivity of the 4 MAbs of cluster III with SakSTAR(E80A,D82A) was maintained in SakSTAR(D82A) but not in SakSTAR(E80A).

7. Absorption of antibodies, elicited in patients by treatment with wild-type SakSTAR

Plasma samples from 16 patients with acute myocardial infarction, obtained several weeks after treatment with SakSTAR (4, 31) were used. The staphylokinase-neutralizing activity in these samples was 20 determined as follows. Increasing concentrations of wild-type or variant SakSTAR (50 μL volumes containing 0.2 to 1000 μ g/mL) were added to a mixture of 300 μ L citrated human plasma and 50 µL buffer or test plasma, immediately followed by addition of 100 μ L of a mixture 25 containing thrombin (50 NIH units/mL) and CaCl, (25 mmol/L). The plasma clot lysis time was measured and plotted against the concentration of SakSTAR moiety. From this curve the concentration of staphylokinase moiety that produced complete clot lysis in 20 min was 30 determined. The neutralizing activity titer was determined as the difference between the test plasma and buffer values and was expressed in μg per mL test plasma. The results of the individual patients have been reported elsewhere (22). For the present invention, three plasma 35 pools were made, one from 10 patients from whom sufficient residual plasma was available, one from three patients that absorbed less than 50% of the antibodies with SakSTAR(K35A,E38A, K74A,E75A,R77A) (Subpool B) and

 $(\hat{f}_{x^{*},T^{*}})$

one from three patients that absorbed >90% of the antibodies with SakSTAR(K35A,E38A,K74A,E75A, R77A) (Subpool C). These plasma pools were diluted (1/30 to 1/200) until their binding to SakSTAR substituted chips in the BIAcore instrument amounted to approximately 2000 RU. From this dilution a calibration curve for antibody binding was constructed using further serial two-fold dilutions. The plasma pools were absorbed for 10 min with 100 nmol/L of the SakSTAR variants, and residual binding to immobilized SakSTAR was determined. Residual binding was expressed in percent of unabsorbed plasma, using the calibration curve.

The results are summarized in Table 1. Whereas wild-type SakSTAR absorbed more than 95% of the binding antibodies from pooled plasma of the 10 patients, incomplete absorption (<60%) was observed with SakSTAR(K74A,E75A,R77A), SakSTAR(K35A,E38A,K74A,E75A,R77A), SakSTAR(K35A,K74A,E75A,R77A), SakSTAR(K35A,K74A,E75A,R77A),

- 20 E38A, K74A, R77A), SakSTAR(K35A, E38A, K74A, E75A),
 SakSTAR(K74A) and SakSTAR(K74A, E75A, R77A, E80A, D82A) but
 absorption was nearly complete with SakSTAR(K35A, E38A),
 SakSTAR(K35A, E38A, E75A, R77A), SakSTAR(E38A, E75A, R77A),
 SakSTAR(E38A, E75A), SakSTAR(K35A, E75A, R77A),
- 25 SakSTAR(K35A,E75A), SakSTAR(E75A), SakSTAR(E80A,D82A), SakSTAR(E80A), SakSTAR(D82A) and SakSTAR(E75A,D82A).

 These results, surprisingly, demonstrate that approximately 40% of the antibodies elicited in patients by treatment with wild-type SakSTAR depend on K74 for
- from 3 patients from which <50% of the antibodies were absorbed with SakSTAR(K35A,E38A,K74A,E75A,R77A) (Subpool B) confirmed the predominant role of K74 for antibody recognition. As expected, absorption with pooled plasma
- 35 from 3 patients from which >95% of the antibodies were absorbed with SakSTAR(K35A,E38A,K74A,E75A,R77A) (Subpool C) was nearly complete with all variants tested.

EXAMPLE 3

Comparative thrombolytic efficacy and immunogenicity of SakSTAR(K74A,E75A,R77A) and SakSTAR(K74A) versus SakSTAR in patients with peripheral arterial occlusion

5 1. <u>Purification of SakSTAR(K74A,E75A,R77A) and</u> <u>SakSTAR(K74A) for use in vivo</u>

A 12 to 24 L culture (in 2 L batches) of the variants SakSTAR(K74A,E75A,R77A), or of SakSTAR(K74A) was grown and IPTG-induced in LB medium supplemented with 100 10 μg/mL ampicillin, pelleted, resuspended, disrupted by sonication and cleared as described above. The compounds were purified by chromatography on a 5 x 20 cm column of SP-Sephadex, a 5 x 10 cm column of Q-Sepharose and/or a 5 x 13 cm column of phenyl-Sepharose using buffer systems 15 described elsewhere (22, 23). The materials were then gel filtered on sterilized Superdex 75 to further reduce their endotoxin content. The SakSTAR variant containing fractions were pooled, the protein concentration was adjusted to 1 mg/mL and the material sterilized by 20 filtration through a 0.22 μm Millipore filter. The methodology used to determine the biological properties of the final material required for use in vivo is described above and elsewhere (22).

25 2. <u>Materials and Methods</u>

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Staphylokinase-neutralizing activity in plasma was determined as described above. Quantitation of antigen-specific IgG and IgM antibodies was performed using enzyme-linked immunosorbent assays in polystyrene microtiter plates essentially as described previously (22). In the IgG assays, dilution curves of affinospecific anti-SakSTAR IgG antibodies were included on each plate. These antibodies were isolated from plasma obtained from 3 patients, after thrombolytic therapy with wild-type SakSTAR, by chromatography on protein A-sepharose and on insolubilized SakSTAR, and elution of bound antibodies with 0.1 mol/L glycine-HCl, pH 2.8. The purity of the IgG preparation was confirmed by sodium

dodecylsulfate polyacrylamide gel electrophoresis. In the IgM assays, titers defined as the plasma dilution giving an absorbancy at 492 nm equivalent to that of a 1/640 dilution of pooled plasma were determined and compared 5 with the titer of baseline samples before treatment (median value 1/410, interquartile range 1/120-1/700).

3. Thrombolytic efficacy

Wild-type SakSTAR or the variants SakSTAR(K74A)

10 or SakSTAR(K74A,E75A,R77A) were administered intra-arterially at or in the proximal end of the occlusive thrombus as a bolus of 2 mg followed by an infusion of 1 mg/hr (reduced overnight in some patients to 0.5 mg/hr) in groups of 6 to 12 patients with

15 angiographically documented occlusion of a peripheral artery or bypass graft of less than 120 days duration. Patients were studied after giving informed consent, and the protocol was approved by the Human Studies Committee of the University of Leuven. Inclusion and exclusion

20 criteria, conjunctive antithrombotic treatment (including continuous intravenous heparin) and the study protocol were essentially as previously described (22).

Relevant baseline characteristics of the individual patients are shown in Table 2. The majority of PAO were at the femoropopliteal level. Two iliac stent and 8 graft occlusions were included. Eight patients presented with incapacitating claudication, 5 with chronic ischemic rest pain, 7 with subacute ischemia and 7 with acute ischemia. One patient (POE) who had 2 years previously been treated with SakSTAR was included in the SakSTAR(K74A) group. This patient was not included in the statistical analyses.

Table 2 also summarizes the individual treatment and outcome. Intra-arterial infusion, at a dose of 35 6.0 to 25 mg and a duration of 4.0 to 23 hrs, induced complete recanalization in 24 patients and partial recanalization in 3. Complementary endovascular procedures (mainly PTA) were performed in 17 patients and

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complementary reconstructive vascular surgery following thrombolysis in 3. No patient underwent major amputation. Early recurrence of thrombosis after the end of the angiographic procedure occurred in 4 patients. Bleeding 5 complications were absent or limited to mild to moderate hematoma formation at the angiographic puncture sites except for 5 patients who required transfusion (data not shown). Intracranial or visceral hemorrhage was not observed. Circulating fibrinogen, plasminogen and 10 α_2 -antiplasmin levels remained essentially unchanged during infusion of the SakSTAR moieties (data not shown), confirming absolute fibrin specificity of staphylokinase at the dosages used. Significant in vivo fibrin digestion occurred as evidenced by elevation of fibrin fragment 15 D-dimer levels. Intra-arterial heparin therapy prolonged aPTT levels to a variable extent (data not shown).

4. Antibody induction

Antibody-related SakSTAR-, SakSTAR(K74A) - and 20 SakSTAR(K74A,E75A,R77A) -neutralizing activity and anti-SakSTAR, anti-SakSTAR(K74A) and anti-SakSTAR(K74A, E75A,R77A) IgG, were low at baseline and during the first week after the infusion (Figure 2). From the second week on, neutralizing activity levels increased to reach 25 median values at 3 to 4 weeks of 20 µg SakSTAR(K74A) and 2 4 µg SakSTAR(K74A) E75A D77A) reutralized remove the second

2.4 μg SakSTAR(K74A,E75A,R77A) neutralized per mL plasma in the patients treated with SakSTAR(K74A) and SakSTAR(K74A,E75A,R77A), respectively, which is significantly lower than the median value of 93 μg

- 30 wild-type SakSTAR neutralized per mL in the patients treated with SakSTAR (p= 0.024 for differences between the three groups by Kruskal-Wallis analysis and p= 0.01 and p= 0.036, respectively, for variants vs wild-type by Mann-Whitney rank sum test). The levels of
- 35 anti-SakSTAR(K74A) and of anti-SakSTAR(K74A,E75A,R77A) IgG increased to median values at 3 to 4 weeks of 270 and 82 μ g/mL plasma in patients treated with SakSTAR(K74A) and SakSTAR(K74A,E75A,R77A) respectively, which is

significantly lower than the median value of 1800 μg anti-SakSTAR per mL plasma in the patients treated with SakSTAR ((p= 0.024 for differences between the three groups by Kruskal-Wallis analysis and p= 0.007 and 0.05, respectively, for variants versus wild-type by Mann-Whitney rank sum test).

The titers of anti-SakSTAR(K74A) and of anti-SakSTAR(K74A,E75A,R77A) IgM increased from median baseline values of 1/460 and 1/410 to median values at 1 10 week of 1/510 and 1/450 in patients treated with SakSTAR(K74A) and SakSTAR(K74A, E75A, R77A), respectively, which was not significantly different from the median values of 1/320 at baseline and 1/640 at week 1 in patients treated with SakSTAR. Corresponding values at 2 15 weeks were 1/590 and 1/550 in patients given SakSTAR(K74A) and SakSTAR(K74A,E75A,R77A), not significantly different from 1/930 with SakSTAR (data not shown). The antibodies induced by treatment with SakSTAR were completely absorbed by SakSTAR but incompletely by 20 SakSTAR(K74A) and by SakSTAR(K74A, E75A, R77A) confirming the immunogenicity of the K74,E75,R77 epitope and the dominant role of K74 in the binding of antibodies directed against this epitope. The antibodies induced by treatment with SakSTAR(K74A) or SakSTAR(K74A,E75A,R77A) 25 were completely absorbed by SakSTAR, by SakSTAR(K74A) and by SakSTAR(K74A, E75A, R77A), indicating that immunization was not due to necepitopes generated by substitution of Lys74 with Ala, but to epitopes different from the K74,E75,R77 epitope.

Thus, this example illustrates that staphylokinase variants with reduced antibody induction but intact thrombolytic potency can be generated. The present experience in 26 patients treated with SakSTAR (n= 9), SakSTAR(K74A) (n= 11) and SakSTAR(K74A,E75A,R77A) (n= 6) combined with previous experience in 14 patients with SakSTAR (n= 7) and SakSTAR(K35A,E38A,K74A,E75A,R77A) (n= 7) (31) and in 24 patients with SakSTAR (32), and with subsequent non-randomized

experience in patients with SakSTAR (n= 30) with SakSTAR(K74A) (n= 12) and with SakSTAR(K74A, E75A, R77A) (n= 7) (data not shown), allows an initial estimation of the prevalence of immunization by intra-arterial 5 treatment with SakSTAR or variants with an altered K74, E75, R77 epitope [SakSTAR(K74A), SakSTAR(K74A, E75A, R77A) and SakSTAR(K35A, E38A, K74A, E75A, R77A)]. Neutralizing activity data after 2 to 4 weeks, available in 70 patients with peripheral arterial occlusion given 10 intra-arterial SakSTAR, revealed that 56 patients (80 percent) had levels > 5 µg compound neutralized per mL plasma. Of the patients given SakSTAR(K74A), SakSTAR(K74A, E75A, R77A) or SakSTAR(K74A, E75A, K74A, E75A, R77A), 27 of the 43 (63 percent) had neutralizing 15 activity levels of > 5 μ g compound per mL plasma. This difference is statistically significant (p= 0.05 by Fisher's exact test) indicating that the K74,E75,R77

20 EXAMPLE 4

Construction, epitope mapping with murine monoclonal antibodies and absorption with pooled plasma of immunized patients, of alanine-substitution mutants of staphylokinase

epitope is a major determinant of antibody induction.

25 1. <u>Introduction</u>

Site-directed mutagenesis was applied to residues other than "charged amino acids" in order to identify i) additional residues belonging to epitopes I and III identified with the panel of murine Mabs and ii) amino acids determining absorption to antiserum from immunized patients. Since functional epitopes generally comprise more than one amino acid residue critical for antibody binding, identification of additional residues in these epitopes could lead to the construction of new combination derivatives displaying a lower antigenic profile, while keeping the specific activity and the temperature stability of wild-type staphylokinase. In this example, the construction and characterization of

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SakSTAR variants in which one or at most two amino acids (adjacent or in close vicinity) were substituted with alanine is described. The mutants described under this example are listed in Table 3. These variants were expressed in <u>E. coli</u>, purified and characterized in terms of specific activity, reactivity with the panel of murine monoclonal antibodies, and absorption of antibodies from plasma of patients treated with wild-type SakSTAR.

10 2. Reagents and Methods

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The source of all reagents used in the present study has previously been reported (22), or is specified below. The template vector for mutagenesis, pMEX602sakB (i.e. pMEX.SakSTAR), has been described elsewhere (23).

- 15 Restriction and modification enzymes were purchased from New England Biolabs (Leusden, The Netherlands), Boehringer Mannheim (Mannheim, Germany) or Pharmacia (Uppsala, Sweden). The enzymatic reactions were performed according to the supplier recommendation. The mutagenic
- 20 oligonucleotides and primers were obtained from Eurogentec (Seraing, Belgium). Plasmid DNA was isolated using a purification kit from Qiagen (Hilden, Germany) or the BIO 101 RPM kit (Vista, CA), as recommended.

 Transformation-competent <u>E. coli</u> cells were prepared by
- the well-known calcium phosphate procedure. Nucleotide sequence determination was performed on double strand plasmid DNA with the dideoxy chain termination method, using the T7 sequencing kit (Pharmacia, Uppsala, Sweden). Polymerase chain reactions (PCR) were performed using Tag
- 30 polymerase from Boehringer Mannheim (Mannheim, Germany) or Vent polymerase (New England Biolabs, Leusden, The Netherlands). The recombinant DNA methods required to construct the variants described in this example are well established (22, 27).

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3. <u>Construction of expression plasmids</u>

The variants SakSTAR(Y17A,F18A), Sak
STAR(F104A), SakSTAR(F111A), SakSTAR(Y9A), SakSTAR(Y91A),

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SakSTAR(Y92A), SakSTAR(I87A), SakSTAR(I106A) and SakSTAR(I120A) were constructed with the Chameleon
Double-Stranded Site-Directed Mutagenesis kit from
Stratagene (La Jolla, USA), using the pMEX.SakSTAR vector
as template, and following instructions of the supplier.
The mutagenic oligonucleotides (not shown) were used in

- The mutagenic oligonucleotides (not shown) were used in combination with the selection-primer LY34 5' CAAAACAGCCGAGCTTCATTCATTCAGC, which destroys the unique HindIII site located 3' to the staphylokinase encoding
- 10 gene in pMEX.SakSTAR and allows to counter-select the non-mutant progeny by HindIII digestion. The deletion of the HindIII site was in most cases correlated with the presence of the desired mutation introduced by the mutagenic oligonucleotide. The variant SakSTAR(I133A),
- 15 was constructed by performing a polymerase chain reaction on the pMEX.SakSTAR plasmid using the primer 818A located at the 5' end of the sakSTAR gene
 - (5' CAGGAAACAGAATTCAGGAG) and the mutagenic primer LY58 (5' TTCAGCATGCTGCAGTTATTTCTTTTCTGCAACAACCTTGG). The
- amplified product (30 cycles: 30 sec at 94°C, 30 sec at 50°C, 30 sec at 72°C) was purified, digested with EcoRI and PstI, and ligated into the corresponding sites of pMEXSakSTAR. The variants SakSTAR(I128A), SakSTAR(L127A) and SakSTAR(N126V) were constructed by performing a
- 25 polymerase chain reaction using the primer 818A located at the 5' end of the sakSTAR gene and mutagenic primers (not shown). The amplified product (30 cycles: 1 sec at 94°C, 1 sec at 50°C, 10 sec at 72°C) was purified, digested with EcoRI and StyI, and ligated into the 30 corresponding sites of pMEXSakSTAR.

The variant SakSTAR(F125A) was constructed by performing two consecutive PCR reactions (30 cycles: : 30 sec at 94°C, 30 sec at 50°C, 30 sec at 72°C). In the first reaction, a fragment of pMEX.SakSTAR was amplified with the primers 818A and a mutagenic primer. This amplified fragment was then used as template in a second PCR reaction with a mutagenic primer in order to further elongate the fragment downstream of the Styl site present

in the sakSTAR gene (corresponding to amino acids 130-131 of SakSTAR). The resulting product was digested with EcoRI and StyI, and ligated into the corresponding sites of pMEXSakSTAR.

- The plasmids encoding all the other variants listed in Table 3 were constructed by direct PCR or by the spliced overlap extension polymerase chain reaction (SOE-PCR)(24) using pMEX.SakSTAR or available plasmids encoding SakSTAR variants as template. Two fragments were
- 10 amplified by PCR (30 cycles: 1 sec at 94°C, 1 sec at 50°C, 10 sec at 72°C), the first one starting from the 5' end (primer 818A) of the staphylokinase gene to the region to be mutagenized (forward primer), the second one from this same region (backward primer) to the 3' end of 15 the gene with primer 818D
- (5' CAAACAGCCAAGCTTCATTCATTCAGC). The forward and backward primers shared an overlap of around 24 bp (primers not shown). The two purified fragments were then assembled together in a second PCR reaction with the
- 20 external primers 818A and 818D (30 cycles: 1 sec at 94°C, 1 sec at 50°C, 10 sec at 72°C). The amplified product from this final reaction was purified, digested with EcoRI and HindIII and ligated into the corresponding site of pMEX.SakSTAR. For each construction, the sequence of
- 25 the variant was confirmed by sequencing the entire SakSTAR coding region.

4. Expression and purification of SakSTAR variants The SakSTAR variants were expressed and

- 30 purified, as described below, from transformed <u>E. coli</u> grown in terrific broth (TB) medium (28). A 2 to 4 mL aliquot of an overnight saturated culture in LB medium was used to inoculate a 1 to 2 L culture in terrific broth supplemented with 100 μ g/mL ampicillin. The culture
- 35 was incubated with vigorous aeration and at 30°C. After about 16 hours incubation, IPTG (200 μ mol/L) was added to the culture to induce expression from the tac promoter. After 3 hours induction, the cells were pelleted by

35 in the table.

centrifugation at 4,000 rpm for 20 min, resuspended in 1/10 volume of 0.01 mol/L phosphate buffer pH 6-6.5 and disrupted by sonication at 0°C. The suspension was centrifuged for 20 min at 20,000 rpm and the supernatant 5 was stored at 4°C or at -20°C until purification. The material was purified essentially as described above (Example 2): cleared cell lysates containing the SakSTAR variants were subjected to chromatography on a 1.6 x 5 cm column of SP-Sephadex, followed by chromatography on a 1.6 x 8 cm column of phenyl-Sepharose. The SakSTAR containing fractions, localized by SDS-gel electrophoresis, were pooled for further analysis.

5. Physicochemical and biochemical analysis

Protein concentrations were determined according to Bradford (29). SDS-PAGE was performed with the Phast SystemTM (Pharmacia, Uppsala, Sweden) using 10-15% gradient gels and Coomassie Brillant blue staining, and the specific activities of SakSTAR solutions were determined with a chromogenic substrate assay carried out in microtiter plates (as described in example 2). The specific activity of the different SakSTAR variants are summarized in Table 3.

25 6. Reactivity of SakSTAR variants with a panel of murine monoclonal antibodies

The methodology used to determine the reactivity of the SakSTAR variants with a panel of murine monoclonal antibodies was described in example 1 above.

30 The results are summarized in Table 3 (the layout of this Table corresponds to the layout of Table 1, as described in example 1). Apparent association constants at least 10-fold lower than those of wild-type staphylokinase were considered as significant and are indicated in bold type

In order to obtain a comprehensive inventory of the properties of Ala-substitution variants of the

SakSTAR molecule, 67 plasmids encoding variants with

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substitution of a single or two adjacent amino acids with Ala were constructed, expressed and purified. Together with the 35 charged residue to Ala-substitution variants previously described (22, and example 2), this analysis covers all residues in SakSTAR except Gly, Ala and Pro, as illustrated in Figure 3. Eight of the variants could not be obtained in purified form, primarily as a result of low expression levels, 11 variants were inactive, 56 had a reduced specific activity, and 27 had a maintained or increased specific activity (≥100 kHU/mg). The yields of purified material from cultures of expressed plasmids were 16 mg/L (median, 10 to 90 percentile range 4 to 41 mg/L). SDS polyacrylamide gel electrophoresis consistently showed one main band with Mr≈ 16,000, usually representing 95% of total protein (not shown).

Substitution of K35, N95, S103 or K135 with Ala yielded variants with specific activities of ≥200 kU/mg. Substitution of W66, Y73 or E75 with Ala reduced the 20 reactivity of the variants with ≥3 antibodies of epitope cluster I, of H43 or V45 with Ala that with 3 antibodies from epitope cluster II and of V32, K35, D82 and K130 with Ala that with ≥3 antibodies of epitope cluster III.

25 7. <u>Absorption of antibodies, elicited in patients by treatment with SakSTAR</u>

For the present example, the three plasma pools, as described in example 2 were used. The methodology used to evaluate the absorption with 30 wild-type staphylokinase and with SakSTAR variants, of antibodies elicited in patients treated with SakSTAR, is described in detail in example 2. The results are summarized in Table 3. Whereas wild-type SakSTAR and most of the variants analyzed in this example absorbed more 35 than 95% of the binding antibodies from pooled plasma of the 10 patients, incomplete absorption (<60%) was observed with SakSTAR(Y73A), and with SakSTAR(K74A). The predominant role of Lys74 for antibody recognition has

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been demonstrated previously (see example 2). The present results indicate that Tyr73 participates to the same major epitope as Lys74, or, alternatively, that substitution at Tyr73 may indirectly induce a structural modification of the "K74-epitope". Absorption with pooled plasma from 3 patients from which >95% of the antibodies were absorbed with SakSTAR(K35A,E38A,K74A,E75A,R77A) (Subpool C, see example 2) was nearly complete with most variants tested.

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EXAMPLE 5

Construction. epitope mapping with murine monoclonal antibodies and absorption with pooled plasma of immunized patients, of staphylokinase variants with substitution of S34, G36 and/or H43

The natural variant Sak42D differs from SakSTAR in three amino acids and corresponds to SakSTAR(S34G, G36R,H43R). Sak42D is characterized by reduced reactivity with some murine antibodies of epitope clusters II and 20 III and a slightly reduced absorption of antibodies from

- plasma of patients treated with SakSTAR (Table 4).

 Mutagenesis of these residues in SakSTAR revealed that
 the reduced reactivity with epitope cluster III and with
 immunized patient plasma could be ascribed to the G36R
- substitution, the H43R substitution mediated the reduced reactivity with epitope cluster II but had no effect on the reactivity with immunized patient plasma, whereas the S34A substitution had no effect. The G36R substitution could be combined with the K74R but not with the K74A
- 30 substitution, without significant reduction of the specific activity (Table 4).

EXAMPLE 6.

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Construction, epitope mapping with murine monoclonal antibodies and absorption with pooled plasma of immunized patients, of staphylokinase variants with substitution of 5 K35, E65, Y73, K74, E80+D82, N95, K130, V132 and/or K135

Based on the results of the alaninesubstitution analysis in example 4, K35, N95 and K135 were selected for further analysis because SakSTAR(K35A), SakSTAR(N95A) and SakSTAR(K135A) had a two-fold increased 10 specific activity, Y73 and K74 because SakSTAR(Y73A) and SakSTAR(K74A) had a markedly reduced reactivity with antibodies from epitope cluster I and diminished absorption of antibodies from plasma of patients immunized by treatment with SakSTAR, and K35, E80+D82, 15 K130 and V132 because SakSTAR(K35A), SakSTAR(E80A,D82A), SakSTAR(K130A) and SakSTAR(V132A) had a reduced reactivity with antibodies from epitope cluster III.

In an effort to maximize the activity/ antigenicity ratio, these amino acids were substituted 20 with other amino acids than Ala. As summarized in Table 5, substitution of K35 with A, E or Q revealed that SakSTAR(K35A) had the most interesting properties, substitution of Y73 with F, H, L, S or W did not rescue the marked reduction in specific activity, and K74 25 confirmed its key role in binding of antibodies from immunized patient plasma, the best specific activity/ antigenicity ratios being obtained with SakSTAR(K740) and SakSTAR(K74R). SakSTAR(E80A, D82A) was preferred over the single residue variants SakSTAR(E80A) or SakSTAR(D82A) 30 because of its somewhat lower reactivity with immunized patient plasma. SakSTAR(N95A) could not be further improved by substitution of N95 with E, G, K or R and it

variants containing K74A or K135R. Finally SakSTAR(K130A) 35 was outperformed in terms of specific activity by SakSTAR(K130T) and SakSTAR(V132A) by SakSTAR(V132R).

was unable to confer its increased specific activity to

EXAMPLE 7

Construction, epitope mapping with murine monoclonal antibodies and absorption with pooled plasma of immunized patients of combination variants of SakSTAR(K130T,K135R)

5 and SakSTAR(E80A,D82A,K130T,K135R) with K35A,G36R,E65X,K74X and selected other amino acids

In the present and the following examples an additional plasma pool was made from 40 patients obtained several weeks after treatment with SakSTAR (Pool 40). The 10 original pool from 10 patients is further identified as Pool 10. The absorption of staphylokinase-specific antibodies was quantified as described above and elsewhere (22).

The SakSTAR(K130T,K135R) variant was taken as a template for additive mutagenesis because of its high specific activity with a moderate reduction of binding to antibodies of epitope cluster III and absorption of antibodies from immunized patient plasma (Table 6). Addition of G36R, K74R, or K74Q or both to the template did not markedly reduce the specific activity, reduced the reactivity with monoclonal antibodies against epitope cluster III (G36R substitution) and decreased the absorption of antibodies from immunized patient plasma (K74R or K74Q substitution). Combination of E65A or E65Q with K74Q in the SakSTAR(K130T,K135R) template reduced the absorption of antibodies from Pool 10 and Pool 40 to around 50 and 60 percent respectively, without markedly

30 K135R) template did not further reduce the antibody absorption from Pool 10 or Pool 40. Surprisingly, the substitution of K136 with A and the addition of K in position 137 resulted in a marked increase in specific activity, as measured in the chromogenic substrate assay.

selected amino acids in the SakSTAR(E65Q, K74Q, K130T,

reducing the specific activity. Addition substitution of

Combination of the SakSTAR(E80A,D82A) and Sak-STAR(K130T,K135R) templates, did not affect the specific activity and had a reduced reactivity with epitope cluster III antibodies (Table 7). Therefore the Sak-

STAR(E80A,D82A,K130T,K135R) template was selected for further mutagenesis. Addition of K74R and even more of K74Q drastically reduced the reactivity with immunized patient plasma. Finally, addition of E65D or of K35A or 5 E65S to the SakSTAR(K74R,E80A,D82A,K130T,K135R) or SakSTAR(K74Q,E80A,D82A,K130T, K135R) templates yielded variants with intact specific activity which only bound ≤45 of the antibodies of pooled immunized patient plasma and less than 15 percent of the subpool reacting for more 10 than 50 percent with the K74,E75,R77 epitope.

EXAMPLE 8

Characterization of selected variants of staphylokinase with intact specific activity and less than 50%

15 adsorption of pooled SakSTAR specific human antibodies elicited in patients by treatment with wild-type SakSTAR

1. <u>Introduction</u>

Twenty three of the variants constructed and characterized in the above examples combined the

20 properties of a residual specific activity of ≥100 kHU/mg and ≤50 percent absorption with the pool of antisera obtained from 10 patients treated with wild-type SakSTAR. The results are summarized in Table 8. Results obtained with Subpool B and Subpool C and with the pool of 40

25 patients treated with wild-type SakSTAR are included. SakSTAR(K74Q,E80A,D82A,K130T,K135R), SakSTAR(K35A,E65D,K74R,E80A,D82A,K130T,K135R), SakSTAR(K35A,E65D,K74Q,E80A,D82A,K130T,K135R) and SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R,K136A,V137K) were selected for further

30 characterization.

2. Fibrinolytic properties of SakSTAR variants in human plasma in vitro

The fibrinolytic and fibrinogenolytic

35 properties of the SakSTAR variants were determined as previously described. Dose- and time-dependent lysis of

125I-fibrin labeled human plasma clots submerged in human plasma was obtained with the selected variants (Table 9).

Spontaneous clot lysis during the experimental period was ≤5% (not shown). Equi-effective concentrations of test compound (causing 50% clot lysis in 2 hrs; C₅₀), determined graphically from plots of clot lysis at 2 hrs 5 versus the concentration of plasminogen activator (not shown), ranged from 0.11 \pm 0.01 to 0.24 \pm 0.04 g/mL at which the residual fibrinogen levels ranges between 92 ± 30 and 97 \pm 30 percent of baseline (Table 9). The concentrations of compound causing 50% fibrinogen 10 degradation in 2 hrs in human plasma in the absence of fibrin were determined graphically from dose-response curves (not shown). These values (mean ± SD of 3 independent experiments) ranged from 14 ± 3.2 to 29 ± 3.1 μg/mL (Table 9). Surprisingly the very high specific 15 activity of SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R, K136A, ∇ 137K) in the chromogenic assay was not associated with an increased thrombolytic potency in a plasma milieu.

- The temperature stability of preparations of SakSTAR(K74Q,E80A,D82A,K130T,K135R), SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R) and SakSTAR(K35A,E65D,K74Q,E80A,D82A,K130T,K135R), dissolved to a concentration of 1.0 mg/mL in 0.15 mol/L NaCl, 0.01 mol/L phosphate buffer, pH 7.5 at various temperatures is illustrated in Fig. 4. At temperatures up to 37°C, all compounds remained fully active for up at least three days. At 56°C and 70°C the three variants were however less stable than wild-type 30 SakSTAR.
 - 4. Pharmacokinetic properties of SakSTAR variants following bolus injection in hamsters

The pharmacokinetic parameters of the 35 disposition of SakSTAR variants from blood were evaluated in groups of 4 hamsters following intravenous bolus injection of 100 μ g/kg SakSTAR variant. SakSTAR-related antigen was assayed using the ELISA described elsewhere.

The ELISA was calibrated against each of the SakSTAR variants to be quantitated. Pharmacokinetic parameters included: initial half-life (in min), $t1/2\alpha = \ln 2/\alpha$; terminal half-life (in min), $t1/2\beta = \ln 2/\beta$; volume of the 5 central (plasma) compartment (in mL), $V_c = dose/(A+B)$; area under the curve (in μ g.min.mL⁻¹), AUC= A/ α + B/ β ; and plasma clearance (in mL.min⁻¹), Clp= dose/AUC (33).

The disposition rate of staphylokinase-related antigen from blood following bolus injection of 100 µg/kg 10 of the selected SakSTAR variants in groups of 4 hamsters could adequately be described by a sum of two exponential terms by graphical curve peeling (results not shown), from which the pharmacokinetic parameters summarized in Table 10 were derived. The pharmacokinetic parameters of the mutants were not markedly different from those of wild type SakSTAR. Initial plasma half-lives (t1/2(a)) ranged between 2.0 and 3.2 min and plasma clearances (Clp) between 1.6 and 4.1 mL/min.

20 EXAMPLE 9

Comparative thrombolytic efficacy and immunogenicity of SakSTAR(K740,E80A,D82A,K130T,K135R) and SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R) versus SakSTAR in patients with peripheral arterial occlusion

25 1. Purification for use in vivo

Eighteen liter cultures (in 2 L batches) of the variants SakSTAR(K74Q,E80A,D82A,K130T, K135R) and SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R) were grown for 20 hours in terrific broth medium (28), supplemented with 100 μg/mL ampicillin and induced with IPTG during the last 3 hours. The cells were pelleted, resuspended in 1/10 volume of 0.01 mol/L phosphate buffer, pH 6.0, disrupted by sonication and cleared by centrifugation. The compounds were purified by chromatography on a 10 x 7 cm column of SP-Sepharose, equilibrated with 0.01 mol/L phosphate buffer, pH 6.0 and eluted with a 1 mol/L NaCl gradient (3 column volumes). The fractions containing SakSTAR variant were pooled, solid NaCl was added to a

concentration of 2.5 mol/L and the material was chromatographed on a 10 x 20 cm column of phenyl-Sepharose followed by stepwise elution with 0.01 mol/L phosphate buffer, pH 6.0. The materials were 5 desalted on a 10 x 45 cm column of Sephadex G25, concentrated by application on a 5 x 10 cm column of SP-Sepharose with stepwise elution with 1.0 mol/L Nacl and finally gel filtered on a 6 x 60 cm column of Superdex 75 equilibrated with 0.15 m Nacl, 0.01 mol/L 10 phosphate buffer, pH 7.5 to further reduce their

10 phosphate buffer, pH 7.5 to further reduce their endotoxin content. The SakSTAR variant containing fractions were pooled, the protein concentration was adjusted to 1 mg/mL and the material sterilized by filtration through a 0.22 m Millipore filter. The

15 methodology used to determine specific activity, endotoxin contamination, bacterial sterility and toxicity in mice is described above and elsewhere (22). The purity of the preparation was evaluated by SDS gel electrophoresis on 10% gels to which 40 g of compound was 20 applied.

Out of culture volumes of 18 liters of SakSTAR variant, 840 mg of SakSTAR(K74Q,E80A, D82A,K130T,K135R) with a specific activity of 140 kHU/mg and 800 mg SakSTAR(E65D, K74R,E80A,D82A,K130T,K135R) with a specific

- 25 activity of 150 were purified. The endotoxin content was <0.1 and 0.26 IU/mg. Gel filtration on HPLC revealed a single main symmetrical peak in the chromatographic range of the column, representing >98% of the eluted material (total area under the curve) (not shown). SDS gel
- 30 electrophoresis of 40 g samples revealed single main components (not shown). Preparations sterilized by filtration proved to be sterile on 3 day testing as described elsewhere (22). Intravenous bolus injection of SakSTAR variants in groups of 5 mice (3 mg/kg body
- 35 weight), did not provoke any acute reaction, nor reduced weight gain within 8 days, in comparison with mice given an equal amount of saline (not shown).

2. Thrombolytic efficacy

Wild-type SakSTAR or the variants SakSTAR(K74Q, E80A, D82A, K130T, K135R) or SakSTAR(E65D, K74R, E80A, D82A, K130T, K135R) were administered intra-arterially at or in 5 the proximal end of the occlusive thrombus as a bolus of 2 mg followed by an infusion of 1 mg/hr (reduced overnight in some patients to 0.5 mg/hr) in groups of 15, 6 and 6 patients respectively with angiographically documented occlusion of a peripheral artery or bypass graft 10 of less than 30 days duration. Patients were studied after giving informed consent, and the protocol was approved by the Human Studies Committee of the University of Leuven. Inclusion and exclusion criteria, conjunctive antithrombotic treatment (including continuous intravenous heparin) and the study protocol were

15 intravenous heparin) and the study protocol were essentially as previously described (22).

Relevant baseline characteristics of the individual patients and results of treatment and outcome are shown in Table 11. Intra-arterial infusion, at a dose 20 of 3.5 to 27 mg and a duration of 2 to 44 hrs, induced complete recanalization in 22 patients and partial recanalization in 5. Complementary endovascular procedures (mainly PTA) were performed in 13 patients and complementary reconstructive vascular surgery following 25 thrombolysis in 5. One patient underwent major amputation. Bleeding complications were usually absent or limited to mild to moderate hematoma formation at the angiographic puncture sites (data not shown). One patient, given wild-type SakSTAR suffered a non-fatal 30 intracranial bleeding, one (BUE) a retroperitoneal hematoma and two (MAN and STRO) a gastro-intestinal bleeding.

Circulating fibrinogen, plasminogen and α_2 -antiplasmin levels remained unchanged during infusion of the SakSTAR moieties (data not shown), reflecting absolute fibrin specificity of these agents at the dosages used (data not shown). Significant in vivo fibrin digestion occurred as evidenced by elevation of fibrin

fragment D-dimer levels. Intra-arterial heparin therapy prolonged aPTT levels to a variable extent (data not shown).

5 3. Antibody induction

Staphylokinase-neutralizing activity in plasma and antigen-specific IgG antibodies were quantitated essentialy as described above and elsewhere (22).

Antibody-related SakSTAR-, SakSTAR(K740,E80A,D82A,

- 10 K130T,K135R) and SakSTAR(E65D,K74R,E80A,D82A,
 K130T,K135R) -neutralizing activity and anti-SakSTAR,
 anti-SakSTAR(K74Q,E80A,D82A,K130T,K135R) and
 anti-SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R) IgG, were
 low at baseline and during the first week after the
- 15 infusion (Figure 5). From the second week on, neutralizing activity levels increased to reach median values at 3 to 4 weeks of 9 μg SakSTAR(K74Q,E80A,D82A, K130T,K135R) and 0.5 μg SakSTAR(E65D,K74R,E80A,D82A, K130T,K135R) neutralized per mL plasma in the patients
- 20 treated with the corresponding moieties, respectively, as compared to median value of 24 μg wild-type SakSTAR neutralized per mL in the 15 patients treated with SakSTAR. The levels of anti-SakSTAR(K74Q,E80A,D82A,K130T,K135R) and of anti-SakSTAR(E65D,K74R,E80A,
- D82A,K130T,K135R) IgG increased to median values at 3 to 4 weeks of 420 and 30 μ g/mL plasma in patients treated with the corresponding moieties, respectively, as compared to a median value of 590 μ g anti-SakSTAR per mL plasma in the patients treated with SakSTAR (Figure 5).
- The prevalence of immunization, defined as neutralizing activities in plasma after 2 to 4 weeks exceeding 5 g/ml was 3 of 6 patients (50 percent) with SakSTAR(K74Q,E80A, D82A,K130T,K135R), 1 of 6 patients (17 percent) with SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R), as compared to
- 35 56 of 70 patients (80 percent) with SakSTAR. This difference is statistically highly significant (p= 0.01 by 2 x 3 Chi square analysis).

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The antibodies induced by treatment with SakSTAR were completely absorbed by SakSTAR but incompletely by SakSTAR(K74Q,E80A,D82A,K130T,K135R) and by SakSTAR(E65D, K74R, E80A, D82A, K130T, K135R) (Table 12).

- 5 Antibodies induced by treatment with Sak-STAR(K74Q, E80A, D82A, K130T, K135R), detectable in 4 of the 6 patients, were completely (≥90 percent) absorbed by SakSTAR, by SakSTAR(K74Q,E80A,D82A,K130T, K135R) and by SakSTAR(E65D, K74R, E80A, D82A, K130T, K135R), indicating that
- 10 immunization was not due to necepitopes generated by substitution of wild-type amino acids. Antibodies induced by treatment with SakSTAR(E65D, K74R, E80A, D82A, K130T,K135R) detectable in one patient (URB) were completely absorbed with SakSTAR(K74Q,E80A,D82A,
- 15 K130T, K135R) and with SakSTAR(E65D, K74Q, E80A, D82A, K130T,K135R) but incompletely (85%) with wild-type SakSTAR, suggesting that a small fraction of the induced antibodies might be directed against a necepitope in the variant used for infusion.

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EXAMPLE 10

Construction and absorption with pooled plasma of immunized patients of combination variants of SakSTAR(E650, K740, K130T, K135R) and other selected amino

25 acids

1. Introduction

In a final round of additive substitution mutagenesis, the SakSTAR(E65Q,K74Q,K130T, K135R) variant was taken as a template because it displayed a high 30 specific activity with a significant reduction of absorption (to 65 percent) of antibodies from pooled immunized patient plasma (Pool 40). The intermediate variants which were relevant for the composition of the finally selected variants are summarized in Table 13. 35 Addition of K35A, D82A and S84A, of T90A, E99D and T101S or of E108A and K109A reduced the antibody absorption to

around 50 percent, whereas the combined addition of D82A,S84A and E108A, K109A reduced it to 41 percent. Substitution of K136A combined with the addition of a Lys at the COOH terminus (-137K) increased the specific activity in a purified system but not in a plasma milieu nor in a hamster pulmonary embolism model (not shown), 5 and further reduced the absorption of antibodies from pooled patient plasma to 30 percent. Finally, addition of the K35A, and T90A,E99D,T101S substitutions to this template yielded a mutant with intact thrombolytic potency which only bound 24 percent of the antibodies of pooled immunized patient plasma.

Based on this analysis, SakSTAR(E65Q,K74Q,D82A, S84A,E108A,K109A,K130T,K135R, K136A,V137K), (SY118), and SakSTAR(K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S, E108A,K109A,K130T,K135R,K136A,V137K), (SY141), were selected for further characterization. In addition, SakSTAR(K35A,E65Q,K74R,D82A,S84A,T90A,E99D,T101S, E108A,K109A,K130T,K135R,K136A,V137K), (SY145) with a Lysin position 74, was constructed and evaluated.

20 2. <u>Pharmacokinetic properties of SakSTAR variants</u> following bolus injection in hamsters

The disposition rate of staphylokinase-related antigen from blood following bolus injection of 100 µg/kg of the selected SakSTAR variants in groups of 4 hamsters could adequately be described by a sum of two exponential terms by graphical curve peeling (results not shown). The pharmacokinetic parameters of the mutants were derived from these plasma disappearance curves not markedly different from those of wild type SakSTAR (results very similar to those of table 10, data not shown).

EXAMPLE 11

Characterization of selected variants derived from SakSTAR(E650, K740, K130T, K135R)

35 1. <u>Fibrinolytic properties of selected SakSTAR variants</u> towards human plasma in vitro

Dose- and time-dependent lysis of ¹²⁵I-fibrin labeled human plasma clots submerged in human plasma was

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obtained with the three selected variants (Table 14). Spontaneous clot lysis during the experimental period was $\leq 5\%$ (not shown). Equi-effective concentrations of test compound (causing 50% clot lysis in 2 hrs; C_{50}),

- 5 determined graphically from plots of clot lysis at 2 hrs versus the concentration of plasminogen activator (not shown), ranged from 0.15 \pm 0.02 to 0.19 \pm 0.01 μ g/ml at which no significant fibrinogen degradation occurred. The concentrations of compound causing 50% fibrinogen
- 10 degradation in 2 hrs in human plasma in the absence of fibrin were determined graphically from dose-response curves (not shown). These values (mean \pm SD of 3 independent experiments) ranged from 7.0 \pm 0.6 to 24 \pm 3.6 μ g/ml (Table 14).

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- 2. Temperature stability of selected SakSTAR variants
 The temperature stability of preparations of
 SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130T,K135R,
 K136A,∇137K), SakSTAR(K35A,E65Q,K74Q,D82A,S84A,T90A,
- 20 E99D,T101S,E108A,K109A,K130T,K135R,K136A,∇137K), and SakSTAR(K35A,E65Q, K74R,D82A,S84A,T90A,E99D,T101S,E108A, K109A,K130T,K135R,K136A,∇137K) dissolved to a concentration of 1.0 mg/ml in 0.15 M NaCl, 0.01 M phosphate buffer, pH 7.5 at various temperatures. At
- 25 temperatures up to 37°C, all compounds remained fully active for up to at least three days. At 56°C and 70°C the variants were generally less stable than wild type SakSTAR (results very similar to those of Figure 4, data not shown).

EXAMPLE 12

Comparative thrombolytic efficacy and immunogenicity of SakSTAR(E650,K740, D82A,S84A,E108A,K109A,K130T,K135R,K136A,V137K), (SY118), SakSTAR(K35A, E650,K740,D82A,S84A,

- 5 T90A, E99D.T101S.E108A, K109A, K130T, K135R, K136A, V∇137K),
 (SY141), and SakSTAR(K35A, E65O, K74R, D82A, S84A, T90A,
 E99D.T101S, E108A, K109A, K130T, K135R, K136A, ∇137K),
 (SY145), in patients with peripheral arterial occlusion
- Large scale purification and conditioning of SakSTAR
 variants for use in vivo

Material was purified to homogeneity out of culture volumes of 18 liters. The endotoxin content was below 2 IU/mg. Gel filtration on HPLC revealed a single main symmetrical peak in the chromatographic range of the

- 15 column, representing >98% of the eluted material (total area under the curve) (not shown). SDS gel electrophoresis of 30 μg samples revealed single main components. Preparations sterilized by filtration proved to be sterile on 3 day testing. Intravenous bolus
- 20 injection of SakSTAR variants in groups of 5 mice (3 mg/kg body weight), did not provoke any acute reaction, nor reduced weight gain within 8 days, in comparison with mice given an equal amount of saline (not shown).

Groups of 6 patients with angiographically
25 documented peripheral arterial occlusion (PAO) were
studied. Relevant baseline characteristics of the
individual patients are shown in Table 15. Table 16
summarizes the individual treatment and outcome.
Intra-arterial infusion, at a dose of 6 to 24 mg and a

- 30 duration of 4 to 29 hrs, induced complete recanalization in most patients. Circulating fibrinogen, plasminogen and α_2 -antiplasmin levels remained essentially unchanged during infusion of the SakSTAR variants (data not shown), reflecting absolute fibrin specificity of these agents at
- 35 the dosages used. Antibody-related SakSTAR(E65Q,K74Q, D82A,S84A,E108A,K109A,K130T,K135R,K136A,∇137K)-, Sak-STAR(K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,

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K109A, K130T, K135R, K136A, ∇ 137K) - and SakSTAR(K35A, E65Q, K74R, D82A, S84A, T90A, E99D, T101S, E108A, K109A, K130T, K135R, K136A, ∇ 137K) -neutralizing activity, were low at baseline and during the first week after the infusion (Table 17).

- 5 From the second week on neutralizing activity levels increased to reach median values at 3 to 4 weeks of 19 μ g SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130T,K135R,K136A, ∇ 137K), (SY118), 0.7 μ g SakSTAR(K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,K135R,
- 10 K136A, ∇ 137K), (SY141), and 4.3 μ g SakSTAR(K35A,E65Q,K74R, D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,K135R, K136A, ∇ 137K), (SY145), neutralized per ml plasma in the patients treated with the respective compounds, which for SY141 and SY145, but not for SY118 is lower than the
- 15 median value of 12 μ g wild type SakSTAR neutralized per ml in 69 patients treated with wild type SakSTAR.

Overt immunization (neutralizing activity at 3 to 4 weeks of 5 g compound per ml plasma) was observed in 56 of 70 patients treated with SakSTAR, in 5 of the 6 patients exposed to SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130T,K135R,K136A,V137K), (SY118), only in 2 of the 6 patients given SakSTAR(K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,K135R,K136A,V137K), (SY141), and in 1 of the 3 patients given SakSTAR(K35A, 25 E65Q,K74R,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,

The results with respect to immunogenicity of the main variants studied in patients are summarized in Table 18. Clearly, variants SakSTAR(E65D,K74R,E80A,D82A, 30 K130T,K135R) and SakSTAR(K35A,E65Q,K74Q,D82A,S84A,T90A, E99D,T101S,E108A,K109A,K130T, K135R,K136A,V137K) have a significantly reduced immunogenicity when compared to the wild type protein.

 $K135R, K136A, \nabla 137K)$, (SY145).

EXAMPLE 13

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Construction, purification and characterization of cysteine-substitution mutants of staphylokinase

1. Introduction

5 Site-directed mutagenesis was applied to substitute exposed amino acids with single cysteine residues in order to construct i) homodimeric forms of staphylokinase, upon formation of an intermolecular disulfide bridge, and ii) polyethylene glycol-conjugated 10 molecules (PEG-derivatives). The aim of this example was twofold: first, the clearance can be reduced by increasing the size of the injected molecule (via dimerization or conjugation with large molecule such as PEG) and second, PEG-derivatives have also been shown to 15 induce a reduced immunoreactivity in animal models (for review, see ref. 34). In both cases, a prolonged half-life in vivo could help to reduce the pharmacological dose of staphylokinase in patients. This reduction could be accompanied with a reduced immunogenic 20 reaction against the thrombolytic agent, thus enhancing its pharmacological activity as a thrombolytic agent.

In this example, the construction and characterization of two SakSTAR variants in which one single amino acid was substituted with cysteine is
25 described. The mutants described under this example are listed in Table 19. These variants were expressed in E. coli, purified and characterized in terms of specific activity, fibrinolytic properties in human plasma in vitro and pharmacokinetic properties following bolus injection in hamsters.

2. Reagents and Methods

The source of all reagents used in the present study has previously been reported (22), or is specified 35 below. The template vector for mutagenesis, pMEX602sakB (i.e. pMEX.SakSTAR), has been described elsewhere (23). Restriction and modification enzymes were purchased from New England Biolabs (Leusden, The Netherlands),

Boehringer Mannheim (Mannheim, Germany) or Pharmacia (Uppsala, Sweden). The enzymatic reactions were performed according to the supplier recommendation. The mutagenic oligonucleotides and primers were obtained from 5 Eurogentec (Seraing, Belgium). Plasmid DNA was isolated

Eurogentec (Seraing, Belgium). Plasmid DNA was isolated using a purification kit from Qiagen (Hilden, Germany), as recommended. Transformation-competent <u>E. coli</u> cells were prepared by the well-known calcium phosphate procedure. Nucleotide sequence determination was

10 performed on double strand plasmid DNA with the dideoxy chain termination method, using the T7 sequencing kit (Pharmacia, Uppsala, Sweden). Polymerase chain reactions (PCR) were performed using Taq polymerase from Boehringer Mannheim (Mannheim, Germany). The recombinant DNA methods 15 required to construct the variants described in this

15 required to construct the variants described in this example are well established (22, 27).

3. Construction of expression plasmids

The variants SakSTAR(K102C) and SakSTAR(K109C),

- 20 were constructed by the spliced overlap extension polymerase chain reaction (SOE-PCR) (24) using pMEX.SakSTAR encoding SakSTAR as template. Two fragments were amplified by PCR (30 cycles: 1 sec at 94°C, 1 sec at 50°C, 10 sec at 72°C), the first one starting from the 5'
- end (primer 818A) of the staphylokinase gene to the region to be mutagenized (forward primer), the second one from this same region (backward primer) to the 3' end of the gene with primer 818D (5' CAAACAGCCAAGCTTCATT-CATTCAGC). The forward and backward primers shared an
- overlap of around 24 bp (for the construction of K102C: TAT GAT AAG AAT TGC AAA AAA GAA GAA (backward) and TTC TTC TTT TTT GCA ATT CTT ATC ATA (forward), for the construction of K109C: AAA AAG AAG AAA CGT GCT CTT TCC CTA (backward) and TAG GGA AAG AGC ACG TTT CTT TTT
- 35 (forward)). The two purified fragments were then assembled together in a second PCR reaction with the external primers 818A and 818D (30 cycles: 1 sec at 94°C, 1 sec at 50°C, 10 sec at 72°C). The amplified product

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from this final reaction was purified, digested with ECORI and HindIII and ligated into the corresponding site of pMEX.SakSTAR. For each construction, the sequence of the variant was confirmed by sequencing the entire coding 5 region.

Expression and purification of SakSTAR variants 4. The SakSTAR variants were expressed and purified, as described below, from transformed E. coli 10 grown in terrific broth (TB) medium (28). A 2 to 4 mL aliquot of an overnight saturated culture in LB medium was used to inoculate a 1 to 2 L culture in terrific broth supplemented with 100 μ g/mL ampicillin. The culture was incubated with vigorous aeration and at 30°C. After 15 about 16 hours incubation, IPTG (200 μ mol/L) was added to the culture to induce expression from the tac promoter. After 3 hours induction, the cells were pelleted by centrifugation at 4,000 rpm for 20 min, resuspended in 1/10 volume of 0.01 mol/L phosphate buffer pH 6-6.5 and 20 disrupted by sonication at 0°C. The suspension was centrifuged for 20 min at 20,000 rpm and the supernatant was stored at 4°C or at -20°C until purification. The material was purified essentially as described above (Example 2): cleared cell lysates containing the SakSTAR 25 variants were subjected to chromatography on a 1.6 x 5 cm column of SP-Sephadex, followed by chromatography on a 1.6 x 8 cm column of phenyl-Sepharose. The SakSTAR containing fractions, localized by SDS-gel electrophoresis, were pooled for further analysis.

5. Biochemical analysis

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Protein concentrations were determined according to Bradford (29). SDS-PAGE was performed with the Phast SystemTM (Pharmacia, Uppsala, Sweden) using 35 10-15% gradient gels and Coomassie Brillant blue staining, and the specific activities of SakSTAR solutions were determined with a chromogenic substrate assay carried out in microtiter plates (as described in

(ii)

example 2). The specific activity of the different SakSTAR variants are summarized in Table 19.

Mutant SakSTAR(K102C) was essentially monomeric as visualized by SDS-PAGE and Coomassie Brillant blue 5 staining. Its specific activity was comparable to that of wild-type staphylokinase. In contrast, SakSTAR(K109C) showed a propensity to form dimers (> 60%). This resulted in a markedly increased specific activity in the plasminogen-coupled chromogenic substrate assay (see 10 Table 19). Upon reduction with dithiothreitol (DTT) (20-fold molar excess during 1.5 hour at 37°C) and alkylation with iodoacetamide (100-fold molar excess during 1 hour at 37°C), the K109C dimer is converted into a stable monomer and its resulting specific activity is 15 within the expected range towards wild-type staphylokinase (Table 19). This result confirms that formation of homodimers is the unique determinant for this large increase in specific activity. Dimeric SakSTAR(K109C) was separated from monomeric 20 SakSTAR(K109C) by chromatography on Source S (Pharmacia) (5 x 50mm). Loading buffer was 10 mM phosphate, pH 6.0 and dimeric SakSTAR(K109C) was eluted by a salt gradient

- (up to 1 M) in the same buffer. The dimeric SakSTAR(K109C) (>95% pure) containing fractions, 25 localized by SDS-gel electrophoresis, were pooled for
- 25 localized by SDS-gel electrophoresis, were pooled for further analysis.

6. <u>Chemical crosslinking of cysteine mutants of SakSTAR</u> <u>with polyethylene glycol</u>

The thiol group of the cysteine mutant SakSTAR(K102C) was targeted for coupling with an activated polyethylene glycol, OPSS-PEG (Shearwater Polymers Europe, Enschede, The Netherlands). OPSS-PEG is a 5 kDa PEG molecule carrying a single activated thiol group at one end that react specifically at slightly alkaline pH with free thiols. Modification of SakSTAR(K102C) was achieved by incubating the molecule (100 μM) with a three-fold excess of SS-PEG in a 5 mM

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phosphate, pH 7.9 solution at room temperature. The extent of the reaction was monitored by following the release of 2-thiopyridone from OPSS-PEG at 412 nm. After reaction (about 15 min), the excess of OPSS-PEG was

5 removed by purifying the derivatized SakSTAR(K102C-PEG) on a 1.6 x 5 cm column of SP-Sephadex as described above (see Example 2). The SakSTAR(K102C-PEG) containing fractions, localized by optical density at 280 nm, were pooled for further analysis. SDS-PAGE analysis and

10 Coomassie blue staining confirmed that PEG crosslinking on SakSTAR(K102C) was quantitative. As shown in Table 19, the specific activity of the PEG-derivative was only marginally affected when compared to that of wild-type staphylokinase.

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7. Fibrinolytic properties of SakSTAR variants in human plasma in vitro

The fibrinolytic and fibrinogenolytic properties of SakSTAR variants were determined as 20 previously described. Dose- and time-dependent lysis of 125 I-fibrin labeled human plasma clots submerged in human plasma was obtained with four molecules: SakSTAR(K109C) as dimer and as monomer (after reduction and alkylation with iodoacetamide), the monomeric SakSTAR(K102C) and the 25 PEG-derivatized SakSTAR(K102C). Spontaneous clot lysis during the experimental period was ≤5% (not shown). Equi-effective concentrations of test compound (causing 50% clot lysis in 2 hrs; C₅₀), determined graphically from plots of clot lysis at 2 hrs versus the concentration of 30 plasminogen activator (not shown), were comparable to that of SakSTAR, for monomeric SakSTAR(K109C) and SakSTAR(K102C) (Table 19). However, it was observed that the C₅₀ for clot lysis by dimeric SakSTAR(K109C) was only 0.12 μ g/ml, which is approximately three-fold lower than 35 for wild-type staphylokinase. In contrast, a C_{50} of 0.60 μ g/ml was measured for SakSTAR(K102C-PEG), which is only two-fold higher than for wild-type staphylokinase. Thus,

dimerization of SakSTAR via disulfide bridges or increasing the size of the molecule via PEG-derivatization does not preclude the fibrinolytic activity of staphylokinase. While a PEG-molecule appears to reduce the diffusion and therefore fibrinolytic potency of the derivatized staphylokinase within a fibrin clot, dimerization of staphylokinase results in a synergistic fibrinolytic effect on human fibrin clots.

10 8. <u>Pharmacokinetic properties of dimeric SakSTAR(K109C)</u> and SakSTAR(K102C-PEG) following bolus injection in hamsters

The pharmacokinetic parameters of the disposition of dimeric SakSTAR(K109C) and SakSTAR(K102C-PEG)

15 from blood were evaluated in groups of 4 hamsters following intravenous bolus injection of 100 μg/kg SakSTAR variant. SakSTAR-related antigen was assayed using the ELISA described elsewhere. The ELISA was calibrated against each of the SakSTAR variants to be quantitated. Pharmacokinetic parameters included: initial half-life (in min), t1/2α= ln2/α; terminal half-life (in min), t1/2β= ln2/β; volume of the central (plasma) compartment (in mL), VC= dose/(A+B); area under the curve (in μg.min.mL⁻¹), AUC= A/α + B/β; and plasma clearance (in mL.min⁻¹), Clp= dose/AUC (32).

The disposition rate of staphylokinase-related antigen from blood following bolus injection of 100 μg/kg of the selected SakSTAR variants in groups of 4 hamsters could adequately be described by a sum of two exponential terms by graphical curve peeling (results not shown), from which the pharmacokinetic parameters t1/2α and Clp, summarized in Table 19 were derived. The pharmacokinetic parameters of dimeric SakSTAR(K109C) and SakSTAR-(K102C-PEG) were markedly different from those of wild type SakSTAR. Initial plasma half-lives (t1/2(α)) were 3.6 and 3.0 min and plasma clearances (Clp) were 0.52 and 0.32 mL/min, for dimeric SakSTAR(K109C) and SakSTAR-(K102C-PEG), respectively. These results may be due to

the increase of the Stokes radius of SakSTAR as a result of the dimerization or crosslinking with PEG. According to size-exclusion chromatography on Superdex50 by HPLC, dimeric SakSTAR(K109C) and SakSTAR(K102C-PEG) have apparent molecular weights of 33 kDa and 40 kDa, respectively.

EXAMPLE 14

maleimide.

Construction, purification and characterization of

10 cysteine-substitution mutants of variants of
staphylokinase with reduced immunogenicity

1. <u>Introduction</u>

Based on the results of example 13, additional polyethylene glycol derivatives of SakSTAR variants were 15 constructed, purified and characterized. The least immunogenic variants SakSTAR(E65D, K74R, E80A, D82A, K130T, K135R), (SY19), and SakSTAR(K35A, E65Q, K74Q, D82A, S84A, T90A, E99D, T101S, E108A, K109A, K1-30T, K135R, K136A, ∇ 137K), (SY141), were used as templates, 20 with the proviso that the COOH-terminus of the latter was reverted to the wild type sequence, S84A was replaced with E80 and K74Q replaced with K74R, yielding Sak-STAR (K35A, E65Q, K74R, E80A, D82A, T90A, E99D, T101S, E108A, K109A,K130T,K135R), (SY161). The introduced cysteine, 25 which functions as acceptor of the polyethylene glycol molecule was located in the amino terminal region (preferably, but not exclusively, the Ser in position number 3 of the mature staphylokinase variant) in order to be released upon activation of staphylokinase (release 30 of the 10 NH2-terminal amino acids); finally polyethylene

The mutants described under this example are

35 listed in Table 20. These variants were expressed in

E.coli, purified and characterized in terms of specific activity, fibrinolytic properties in human plasma in vitro, pharmacokinetic properties following bolus

to 20,000) were used, substituted with either OPSS or

glycol molecules of different molecular weights (M 5,000

injection in hamsters, thrombolytic properties following bolus injection in a hamster pulmonary embolism model, and absorption of antibodies from pooled immunized patient plasma (Pool 40).

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2. Reagents and Methods

The source of all reagents used in the present study has previously been reported (22), or is specified below. The template vector for mutagenesis, pMEX602sakB 10 (i.e. pMEX.SakSTAR), has been described elsewhere (23). Restriction and modification enzymes were purchased from New England Biolabs (Leusden, The Netherlands), Boehringer Mannheim (Mannheim, Germany) or Pharmacia (Uppsala, Sweden). The enzymatic reactions were performed 15 according to the supplier recommendation. The mutagenic oligonucleotides and primers were obtained from Eurogentec (Seraing, Belgium). Plasmid DNA was isolated using a purification kit from Qiagen (Hilden, Germany), as recommended. Transformation-competent E. coli cells 20 were prepared by the well-known calcium phosphate procedure. Nucleotide sequence determination was performed on double strand plasmid DNA with the dideoxy chain termination method, using the T7 sequencing kit (Pharmacia, Uppsala, Sweden). Polymerase chain reactions 25 (PCR) were performed using Taq polymerase from Boehringer Mannheim (Mannheim, Germany). The recombinant DNA methods required to construct the variants described in this example are well established (22, 27).

30 3. Construction of expression plasmids

The variants SakSTAR(S3C,E65D,K74R,E80A,D82A,K130T,K135R), (SY19(S3C)), SakSTAR(S2C,S3C,E65D,K74R,E80A,D82A,K130T,K135R), (SY19(2SC,3SC)), SakSTAR(S3C,K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,K135R,K136A,V137K), (SY141(S3C)), SakSTAR(S2C,S3C,K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,-K109A,K130T,K135R,K136A,V137K), (SY141(S2C,S3C)), SakSTAR(S3C,K35A,E65Q,K74Q,E80A,D82A,T90A,E99D,T101S,E108A,

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K109A, K13OT, K135R), (SY160(S3C)) and SakSTAR(S3C, K35A, E65Q, K74R, E80A, D82A, T90A, E99D, T101S, E108A, K109A, K130T, K135R), (SY161(S3C)), were constructed by the spliced overlap extension polymerase chain reaction 5 (SOE-PCR) (24) using pMEX.SakSTAR encoding SakSTAR as template, two fragments were amplified by PCR (30 cycles: 1 sec at 94°C, 1 sec at 50°C, 10 sec at 72°C), the first one starting from the 5' end (primer 818A) of the staphylokinase gene to the region to be mutagenized 10 (forward primer), the second one from this same region (backward primer) to the 3' end of the gene with primer 818D (5' CAAACAGCCAAGCTTCATTCATTCAGC). The forward and backward primers shared an overlap of around 24 bp. The two purified fragments were then assembled together in a 15 second PCR reaction with the external primers 818A and 818D (30 cycles: 1 sec at 94°C, 1 sec at 50°C, 10 sec at 72°C). The amplified product from this final reaction was purified, digested with EcoRI and HindIII and ligated into the corresponding site of pMEX.SakSTAR. For each 20 construction, the sequence of the variant was confirmed by sequencing the entire SakSTAR coding region.

4. Expression and purification of SakSTAR variants The SakSTAR variants were expressed and

purified, as described below, from transformed E. coligion of an overnight saturated culture in LB medium was used to inoculate a 1 to 2 L culture in terrific broth supplemented with 100 μg/mL ampicillin. The culture was incubated with vigorous aeration and at 30°C. After about 16 hours incubation, IPTG (200 μmol/L) was added to the culture to induce expression from the tac promoter. After 3 hours induction, the cells were pelleted by centrifugation at 4,000 rpm for 20 min, resuspended in 1/10 volume of 0.01 mol/L phosphate buffer pH 6-6.5 and disrupted by sonication at 0°C. The suspension was centrifuged for 20 min at 20,000 rpm and the supernatant was stored at 4°C or at -20°C until purification. The

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material was purified essentially as described above (Example 2): cleared cell lysates containing the SakSTAR variants were subjected to chromatography on a 1.6 x 5 cm column of SP-Sephadex, followed by chromatography on a 1.6 x 8 cm column of phenyl-Sepharose. The SakSTAR containing fractions, localized by SDS-gel electrophoresis, were pooled for further analysis.

5. Biochemical analysis

Protein concentrations were determined according to Bradford (29). SDS-PAGE was performed with the Phast System™ (Pharmacia, Uppsala, Sweden) using 10-15% gradient gels and Coomassie Brillant blue staining, and the specific activities of SakSTAR solutions were determined with a chromogenic substrate assay carried out in microtiter plates (as described in example 2).

6. Chemical crosslinking of cysteine mutants of SakSTAR with polyethylene glycol

The thiol group of the cysteine mutants was targeted for coupling with an activated polyethylene glycol, either OPSS-PEG or MAL-PEG (Shearwater Polymers Europe, Enschede, The Netherlands). OPSS-PEG is a 5 kDa 25 PEG molecule carrying a single activated thiol group at one end that reacts specifically at slightly alkaline pH with free thiols. MAL-PEG is a 5 kDa, 10 kDa or 20 kDa molecule carrying a maleimide group that reacts specifically with thiol groups under mild conditions in the 30 presence of other functional groups. Modification of the variants was achieved by incubating the molecule (100 μ M) with a three-fold excess of OPSS-PEG or MAL-PEG in a 5 mM phosphate, pH 7.9 solution at room temperature. After reaction (about 15 min), the excess of OPSS-PEG or 35 MAL-PEG was removed by purifying the derivatized SakSTAR variant on a 1.6 x 5 cm column of SP-Sephadex as described above (see Example 2). The "pegylated" SakSTAR variant containing fractions, localized by optical densii e

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ty at 280 nm, were pooled for further analysis. SDS-PAGE analysis and Coomassie blue staining confirmed that PEG crosslinking was quantitative. As shown in Table 20, the specific activities of the PEG-derivatives were only marginally affected when compared to that of wild-type staphylokinase.

7. Fibrinolytic properties of SakSTAR variants in human plasma in vitro

- 10 The fibrinolytic and fibrinogenolytic properties of SakSTAR variants were determined as previously described. Dose- and time-dependent lysis of ¹²⁵I-fibrin labeled human plasma clots submerged in human plasma was obtained with all molecules tested.
- 15 Equi-effective concentrations of test compound (causing 50% clot lysis in 2 hrs; C₅₀), determined graphically from plots of clot lysis at 2 hrs versus the concentration of plasminogen activator (not shown), were comparable to or only slightly lower than that of SakSTAR (Table 20). The
- 20 C_{50} for clot lysis by variants derivatized with P20 (PEG with \underline{M}_r 20 kDa) was about twice as high as the non-derivatized variants. Thus increasing the size of the molecule via PEG-derivatization does not markedly affect the fibrinolytic activity of staphylokinase. The
- 25 PEG-molecules appear to reduce the diffusion and therefore fibrinolytic potency of the derivatized staphylokinase within a fibrin clot, but this appears to be less pronounced with variants substituted in their NH₂-terminal region which is released during processing of staphylokinase than with variants substituted in the
 - 8. <u>Pharmacokinetic properties of SakSTAR variants</u> <u>chemically modified with polyethylene glycol</u>

core of the molecule (cfr. Tables 19 and 20).

following bolus injection in hamsters

The pharmacokinetic parameters of the disposition of the pegylated variants from blood were evaluated in groups of 4 hamsters following intravenous

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bolus injection of 100 μ g/kg SakSTAR variant. SakSTAR-related antigen was assayed using the ELISA described elsewhere. The ELISA was calibrated against each of the SakSTAR variants to be quantitated.

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5 Pharmacokinetic parameters included: initial half-life (in min), t1/2α = ln2/α; terminal half-life (in min), t1/2β = ln2/β; volume of the central (plasma) compartment (in mL), VC= dose/(A+B); area under the curve (in μg.min.mL⁻¹), AUC= A/α + B/β; and plasma clearance (in mL.min⁻¹), Clp= dose/AUC (32).

The disposition rate of staphylokinase-related antigen from blood following bolus injection of 100 μg/kg of the selected SakSTAR variants in groups of 4 hamsters could adequately be described by a sum of two exponential terms by graphical curve peeling (results not shown), from which the plasma clearances Clp, summarized in Table 20 were derived. The clearances of pegylated variants were markedly different from those of wild type SakSTAR and were inversely proportional to the molecular weight of the PEG molecules, with an average reduction of 5-fold with PEG 5 kDa, 10-fold with PEG 10 kDa and 30-fold with PEG 20 kDa. These results may be due to the increase of the Stokes radius of SakSTAR as a result of crosslinking with PEG.

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EXAMPLE 15

Comparative thrombolytic efficacy and clearance of Sak-STAR(S3C-P20,E65D,K74R, E80A,D82A,K130T,K135R).
(SY19(S3C-P20)), in two patients with acute myocardial

30 <u>infarction</u>

Large scale purification and conditioning of the SakSTAR variant for use in vivo

Material was purified to homogeneity out of culture volumes of 18 liters. The endotoxin content was 35 below 1 IU/mg. Gel filtration on HPLC revealed a single main symmetrical peak in the chromatographic range of the column, representing >98% of the eluted material (total area under the curve) (not shown). SDS gel

electrophoresis of a 30 μg sample revealed single main component. The preparation sterilized by filtration proved to be sterile on 3 day testing as described in methods. Intravenous bolus injection of the SakSTAR
5 variant in 5 mice (3 mg/kg body weight), did not provoke any acute reaction, nor reduced weight gain within 8 days, in comparison with mice given an equal amount of saline (not shown).

Two patients with acute myocardial infarction

10 were given a bolus injection of 5 mg SY19(S3C-P20). These
patients had a complete recanalization of the occluded
infarct-related artery as determined by coronary
angiography at 90 min after the bolus injection. The
material was cleared from the plasma with an initial

15 half-life of 3 to 4 hours, as compared to 4 to 6 minutes
for wild-type SakSTAR. These data confirm that pegylated
variants of SakSTAR may be useful for thrombolytic
therapy by single bolus injection at a reduced dose.

20 CONCLUSION

(5)

In summary, the present invention shows that staphylokinase variants with markedly reduced antibody induction but intact thrombolytic potency can be generated. This observation constitutes the first case in 25 which a heterologous protein, with the use of protein engineering techniques, is rendered significantly less immunogenic in man without reducing its biological activity. In addition, the present invention shows that selective chemical modification of staphylokinase or its 30 variants with polyethylene glycol of varying molecular weights is feasible, resulting in a reduction of the plasma clearance proportional to the molecular weight. In the preferred embodiment an amino acid in the NH,-terminal region of staphylokinase, the portion that is removed by 35 processing, is substituted with Cys and the introduced thiol group is chemically modified with OPSS-PEG or MAL-PEG. This results in homogeneous products which, upon single intravenous bolus injection in experimental

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animals and in patients have a maintained thrombolytic

potency at markedly reduced doses.

Alanine-to-wild-type" reversal variants of "charged-cluster-to-alanine" mutants of SakSTAR: Association constants (KA x 107 mol/L-1) for the binding to insolubilized murine monoclonal antibodies (Mabs), and absorption (percent) of antibodies of immunized patient plasma Table 1:

Variant	Exp.	Spec. Act.		ľ	Epitope 1		-		murine MAbs	44×			ľ	Epitope	L		Saks	SaksTAR patient	Dasma
	(mg/L)	(kU/mg)	1301	26A2	L	2817	3610 18F12	THE LANS	L_	3282	FEB		23E1	ğ	7307	1710	8		Subpool C
SakSTAR		06.1	73	F	ľ	8	86	7	۵	F	7.	6.4	÷	k	£	9,0	32	į,	8
SakSTAR(K35A.E38A)		66	2	22	4.2	1.9	=	2	2	2	2.2	₹ 7	9.1	9.1	0.1	9:	93	5	3
Saestar(K14A,E75A,R77A)		110	=	6 0	6	6.1 6.	150	1	78	=	33	2.4	=	0.4	2.1	6.0	55	\$	8
SekSTAR(K35A.E38A,K74A.E75A.R77A)		20	=	&	.6 .1	.e.		8	92	2	2.0	6.1	9.1		<u>.</u>	7:1	ĸ	=	8
SakSTAR(E38A,K74A,E75A,R77A)		43	=	9.1	-Q.2	.i.	- - - - -	8	92	2	7.1	8	3.2	3.7	9.	=	\$	\$	93
SakSTAR(K35A, K74A,E75A,R77A)		28	9.2	.	0.15	.e.	1 32	=	53	89.	23	<u>ē</u>	9 2	8 .	9 .	8.0	\$	\$	8
SakSTAR(K3SA.E38A.E7SA.R77A)		2	Ξ	0.3	0.1	0.2 <0.1	25	9.6	2	7.3	9.	8.	9.1	9.1	0.53	29:0	33	81	8
SakSTAR(K35A.E38A.K74A.R77A)		4	89.	2.9	40.1 2.	2.0 0.33	9 =	33	3	9	2.0	9.1	9	9.	0.63	0.74	38	8	93
SINSTAR(K3SA.E38A.K74A.E75A)		2	2	6.5	 	6.1 6.1	281	4	37	51	9:	6.	9.1	ē	7	0.45	#		85
SakSTAR(E38A.E75A.R77A)		88	=	9.0	0.15 0.	4 0.3	٤	13	2	2	2.0	ę	2.6	4.7	Ξ	0.81	26	80	5 (
SakSTAR(E38A,E75A)		*	9	0.3	6.1	<0.1 0.9	%	=		8.9	2.0	9.	2	8.	<u>:</u>	9.	5	06	8
SakSTAR(K35A.E75A.R77A)		89	9.2	6	6.1	.6. .6.	8	7.0	2	=	3.3	9	2	9.	0.8	=	80 80	8	. 8
Sakstar(K35A,E75A)		150	11	0.12	d.1 0.	0.16 0.14	\$	7.2	=	9.3	4.2	9.1	8 :	ē.	<u>*</u>		8	93	88
SakSTAR(K74A)		901	13	9.7	0.17 4.4	4 2.1	S	15	33	4	3.6	2.9	4	4.9	3.4		\$	\$	88
SakSTAR(E75A)		140	=	2	.0°.	<u>6</u>	*	8.5	7	2	3.4	\$	<u>e</u>	8.0	17		8	93	8
SakSTAR(K74A.E75A.R77A.E80A.D82A)		80	*	9.1	6.	6	8	6	33	6	3.7	9.	6	9.1	9.1	7.7	\$	29	88
Sakstar(E80A,D82A)		130	1.3	2	9	5.9	۶	9.	8.4	7.8	<u>•:</u>	9.	å.			1	2	2	33
Sakstar(E80A)		8	13	=	3 7.9	9	35	7.4	1	8.6	2.1	6	9	3.6			æ	93	88
SakSTAR(D82A)		8	11	12	.7	=	=	7.8	11	2	2.7		0.18	9.1	6.1		95	93	8
Sakstar(E75A.D82A)		170	22	<u>5</u>	3.1 6.6	6 7.2	\$	8	2	7	6.4	0.17	0.7	0.5	1 170	<u></u>	8	. 86	88
							_								-		-		1

Apparent association constants ≥ 10-fold lower than those of wild-type SakSTAR are represented in bold type; Spec. Act. ≥ 100.000 HU/mg represented in bold type; ≤60% absorption represented in bold type.

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Table 2:
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SakSTAR MEE			(days)	(cm)	Inrombolysis	thrombolytic agent (mg)	of infusion (hrs)	
Σ Σππ Σ ΣΣ Σ ,	Rest pain	Len SFA	30	80	complete	7.0	\$.0	· PTA
Σππ Σ ΣΣ Σ ,	Claudication	Left IA (stent)	4	18	complete	. 6.5	\$	PTA + stent
- Err 2 22 2	Claudication	Dioh: CEA	Ç	4	a) al amon	,	Š	į
ا۔ ت ک کک ک '	Deet pain	Les ET and	R 2	> \$	complete	. <u>.</u>		YIA
Σ ΣΣ Σ -	Acute	Left brachial and	2 2	3 ~	complete	<u>•</u> •	2 1	PTA + steni
, 2		radial artery				:		
ΣΣ Σ ,	Claudication	Right SFA (popliteal	%	13	complete	0.9	4.0	PTA + femoropopliteal bypass
, Z Z -	Acute	I of PA	-	ç	e le le le le		7.0	graff
Σ -	Acute	Left EIA (stent)	4	2 2	complete	89	5. 4	(amputation teft digit V)
-	Subacute	Right FP graft	e	45	complete	0	0.9	3
			17±5.6	21 ± 5.8		9.7 ± 1.7	9.1 ± 2.7	
LIE M 70	Subscute	Richt FF oraft	9	48	applance	=	c	Š
	Claudication	Right SFA	, c	2	complete	: 2	? =	474
<u>.</u>	Claudication	Right PA graft	2 2	۲.	narrial	: =	2 ¥	PTA
Ŀ	Claudication	Right SFA	>120	• •	complete	0.6	2.0	ATQ
	Acute	Right IF graft	2	*	complete	<u>~</u>	9	Surpical graft revision
	Acute	Right IF and FP graft	_	63	complete	2	2	PTA
: -	Rest pain	Right TF trunc	0.6	38	partial	8 2	71	•
L .	Rest pain	Left AF graft	23	78	сотрієє	<u></u>	7	•
Σ:	Subacute	Right TF trunc	7	R	partial	0.9	4.0	n-PA, surgical graft lengthening
VBE M 39	Subacute	Right BA (embolism)	70	38	complete	82	23	Stent right SC artery. first rib
SME F 50	Subacute	TF trunc	88	32	complete	21	61	None
W 67	Subacute	Right PA	4	23	complete	91	22	•
5±30		•	23 ± 9.2	35 ± 6.4		15±1.2	16±1.9	
SEKSTAR(K74A,E75A,R77A)								
∟ :	Acute	Right BA and UA	0.3	~	complete	7	2	
MAE M 74	Rest pain	Len SFA	2	S	complete	0.6	7.0	PTA
	Claudication	Right IA and FA	4	28	complete	22	23	PTA + stent
W 68	Claudication	Left SFA	8	12	complete	. 0.6	7.0	PTA
Z Z	Subacute	Len SFA	4	•	complete	0.6	7.0	PTA
Z 20	Acute	Right FT graft		42	complete	0.6	7.0	MA ATA
Mean ± SEM 65±3.3			22±14	24±7.8		13 ± 2.6	11 ± 2.6	

Alanine-substitution variants of SukSTAR: Association constants (K, x 10°mol/L.¹) for binding to insolubilized murine monoclonal antibodies (Mab) and absorption (percent) of antibodies of immunized patient plasma 2 2 2 2 2 2 2 2 2 1 2 8 20 0.75 3.8 형 é 夸 e. 6 5.0 20 5 2 50 £ 9 77 100 20 8.0 충 3. 20 5.4 Ç 52 33 3.6 2 • 3.6 6 é 7.6 3.4 Ξ 9.6 Ş wi. 53 7 Ξ 2 章 5.6 6 9 ~ ~ 2 뤽 Ξ 2.7 2 56 3.6 2 0 7 33 3 2 2 73 2 2 <u>*</u> 2 ~ €. 2 2 2 룬 9.0 5. 5. = 5 7.5 0.7 6.2 5. 2 2 9.2 6, 2 9.0 2 2 2 2 2 7.7 9. 7, 17GH 26AZ 2 9.6 2 Exp. Spec. Act. (mg/L) (kU/mg) 8 2 8 ŋ \$ 9 5 5 3 SASTARIKI I A.DIJA.DI4A) ASTAR(S34G,G36R,H43R) SakSTARID!!A.K!!SA! akSTAR(Y17A.F18A) LAKSTAR(N28A,V29A) akSTAR(KBA,K10A) STARIE19A.P20A BASTAR(DSA,K6A) SASTAR(DI3A) ASTAR(M26A) SakSTAR(F4A) akSTAR(Y9A) akSTAR(D14A) **ILSTARISIBA** ASTAR(T21A) akSTAR(P23A) UNSTARIY24A) #STAR(L25A) SASTARIV27A) LESTAR(N2BA) SASTAR(T30A) SatSTARIV32A) SHSTARISHAD Table 3: Varian

Table 3 - cont'd: Alanine-substitution variants of SakSTAR: Association constants (K, x 10'mol/L') for binding to insolubilized murine monocional antibodies (Mab) and

absorption (percent) of an	rcent) o	f antibodi		titis of Sand I And Association constant tilbodies of immunized patient plasma	unized	pati	ent pie	STA	Y (1)					9				of immunized patient plasma		
			<u>ا</u>							É	murine MAbs	E I					\prod			
Vertent	(Beg.	Spec. Act.		17011 28	26A7 30A2 2B12		3010	THEFT	36	Market Chart	Ė		TIE!	Ē			IVIO	r IooL	Subpool B	Subpool
SAESTAR(ROSA)		2	_		F	=	F	F	F	F	L	Т	FE S	ı	,	-	80	5	22	88
SakSTAR(KJSA,EJ8A)		6		15 22	4.2	=	7.9	<u>=</u>	2	2	~	77	1.	ě	Ą	0.1	0.	8	5	z
SabSTAR(Q36A)	<u> </u>	2		8.9.8	5.1	5.7	2,	8	7	=	9.7	<u> </u>	Ŧ	Ŧ	6.0	5.0	<u>e</u>	*	8	8 2
SakSTAR(N17A)	9	2	<u> </u>		3.0	2	=	8	7	=	2	2.9	<u>.</u>	53	2.5	3.6	8	8	8	8
Sakstariljoa.L40a)	-	৯	<u> </u>	55	3	3.1	0.	3	92	63	=	2.7	7.	3.4	3.2	7.	6.0	8	8	8
SekSTAR(\$41A,P42A)	*	=		22	₹	=	11	<u>=</u>	3.0	1.9	=	2.7	3.2	2	8.	3.6	Ξ	8	88	86
SatSTAR(H43A)	ĸ	8	=	28	9.7	=	1.6	ş	6	41.9	9.1	<u>5</u>	5.0	ສ	7.8	27	9:	8	8	2
Sakstar(H1)A,Y41A)	=	v		22	3.7	-	2	ŧ	ş	동 -	3	30	ม	=	23	7.	-6	8	2	8
SakSTAR(V45A)		ŋ	<u> </u>	5.6	-	7	63	2	0.2	1.7		- -	- I	CB	Ξ	2.8	9.	5	~	8
Sakstar(E46A.K50A)	밀																			
SakSTAR(F47A)		্হ	ę	0.1 4.0	0.	3.9	3.4	5.7	1.7	2.8	8.5	٠.	6.0	80	3.0	3.0	6.0	8	2	93
SakSTAR(149A)	~	7	1.7	" "	7.8	2	Ħ	×	3	9	6.2	=	02	5.7	07	1.7	9.0	8	88	8
SakSTAR(KSOA)	5	7	₹	5	2.9	7.8		ş	2	2	0 77		2.8	9.4	0.4	23	9.0	8	z	86
SakSTAR(T33A,T34A)	=	\$	\$	61	1.1	7.6	7.8	<u> </u>	6.3	2		<u> </u>	61.	5.1	2	0.1	9.0	2	z	88
SakSTAR(L.55A)	3																			
SakSTAR(T36A)	=	8	<u>.</u> 2	5	3.2	2	=	8	5.3	2	=	- C 02	35	1.9	1.1	7	12	z	26	8
SakSTAR(KS7A,ES8A,KS9A)		8	<u>=</u>	8.7	9	ני	11	9	±	6.7 5.	3.6	0.52	0.36	:	0.42	0.1	Ξ			
Sabstar(160A)	=	8		.2	2.9	=	=	æ		. 22	2.7	<u>ه</u>	0.7	5.8	62	1.7	2	88	8	8
SakSTAR(E61A,E65A)		8	19.5	5 × 50	8.8	=	2	ž	914	6.6	>7.2 4	6.0	50	4.6	2.0	5.9	2			
SukSTARIY61A.Y63A)	Σ.	♡	₹		0.3	2.1	6:	_=	2	3.1	-	<u></u>	9.0	9.3	3.6	3.8	0.7	2	ä	93
SakSTAR(Y63A)	_=	হ	₹	# I	1.1	9.6	Ξ	2	2	25	25	- 7 	77	53	Ç	0.1	7.	&	83	88
SetSTAR(V64A)	<u> </u>	\$	=	91	2.9	6.3	7.8	2	2	21 23		- <u>-</u> -	1.6	7.6	3.6	3. 8.	0.1	a	8	8
Saltar(E65A)	×		<u>.s</u>	8	7	=	0.7	<u> </u>	3.6	9.7	. 6:	1.8		4.7	3.0	5.8	0.97	8	8	
Salstareesa. D69a)	<u></u> -	ঢ়																		
SahSTAR(W66A)	29	ซ	₹	<u>6</u>	鱼	章	₹	2	\$	5.7	ъ К	33 20		9	<u>.</u>	=	8.0	28	18	26
SakSTARILGBA)	2	2	=	=	3.5	8 .5	63	£	8.7	5	¥ \$1	4.0		53	3.6	7	ŧ	8	8	2
Salstar(T)!AI	빌											_								
_	_	_	_									-					-			٠

Table 3 - cont'd: A	Alanine-substitution variants of SakSTAR: Association constants (K _A x 10'mol/L'¹) for binding to insolubilized murine monocional antibodies (Mab) and absorption (percent) of antibodies of immunized patient plasma	on varia	ints of	SakST es of in	AR: /	Associ	ation	Const	ants (KAX	10°mol	T.') f	or bin	ling to	Insolub	llized m	urine n	sonocion	al antibodi	es (Mab) and
Variant	- En	Spec		Н	o doing	Talent Talent					munine MAbs			ľ						
	(mg/L)	L) (kU/mg)		17511 28,	F	26A2 30A2 2B12	2,510	THEFT	1	28HT 37B	L.	110	THE .	381	40C8 24C		B.Y	7	SakSTAR patient plasma	різата
Saks (AR(Y7,(A)	<u>e.</u>	0	-	3	ı	ત્વા ત્વા		=	þ	F		Т			5	Ŀ	2.60	.1	a loodens	Supposite S
SubSTAR(Y7:1A.K74A)	77	ŋ	=	를	<u>6</u>	.±.	<u>6</u>	<u>•</u>	6.3	23	9.9 3.2		2.7	2	0,	9.	=	ţ	*	83
SakSTAR(K74A)	<u>&</u>	\$		1.2.7	6.2	11	Ξ	-	5.2	=	7.6 2.2		2.0	89	5	8.1	6	3	. S	. 5
SakSTARIK74A.E75A.R77A)		3	2.	<u>é</u>	<u>6</u>	6.	₹	8	0.7	=	33	<u> </u>	=	*	ਦ	8.0	=	. 22	2	: 5
SakSTAR(K74A,R77A)	<u> </u>	-	3.5	=	0.2	<u>.</u>	3	2	7.7	2	2.1 1.8	_ <u>=</u> _	7	23	::	~			\$ \$: \$
SatSTAR(E75A)		<u> </u>	=	2	Ġ.	- G	Ą	\$	2	=	3.4			∞	20	~	17	. ×	: :	: 8
· SakSTAR(F76A)		8	2	9.6	1.0	2,3	3.9	_=	79	2	13		_	5.9	7.	~	9	: a	: 8	: :
SakSTAR(V78A,V79A)	<u> </u>	89	2	2	0.4	5	1	=		**	1.2	<u> </u>	6		é.	20	-	. 5	. 16	. 2
SakSTAR(E80A)		3	2	=	33	7.9	2	25	7.	2	8.6 2.1	6.	22		3.6	ਵ	-	. 2	: 5	. 56
SalsTAR(EBOA, D82A)		옷	2,2	2	2.1	6.5	\$.9	۶		1,4	7.8 1.9	<u>₹</u>	ž. Ą	_	ą.	를	3	2	: 2	: 8
SakSTAR(LBIA)	22	28	~	2	9.	\$	=	2	=	-	17 3.9		22		17	6	-	.		: 8
SakSTAR(D82A)		<u> </u>	=	~	*	1.3	=	_5	7.8		12 23	<u> </u>	_		_	· =	: :	8 8	: 6	: *
SalsTAR(DEZA,SBAA)	22	5	<u>.</u>	=	2.6	.	5.8	_2	3.6	-	= 5	ਵ	_	_		ਵ	9	: 5	: 5	. 2
SakSTAR(S84A)	12/26	<u>£</u>	0.8	2	3.8	9.6	9	_8	3	=	36	7.7	9.		3.0	3.5	20	. 2.	. .	: S
SakSTAR(K86A.E88A)		2	[7.2	3	33	9.0	9;	5.7	7. 6.4	. 15	\$	6.	3.4		0.80	<u>6:</u>	0.0			
SukSTAR(187A)	<u></u>	8	6.7	2	2.8	8.6	ē	9	3.6	7.4	2.7	_=	7.8		3.4	.	9	\$6	8	2
SakSTAR(V89A)	20	<u>&</u>	4.	=	2.6	9.6	77	*	1.2.7		3.0			~		<u> </u>	0.83	. 2	: x	: 2
Sakstar(T90A)	78	2	9.0	2	6.0	3.7		2	4.8	5. 2.			9.6	2,6				. 56	. x	. 2
SabSTAR(Y91A)	<u>.</u> ~	\$	9.0	2	3.0	7.0	2	_=	8.2 16	3	7		3.7	-	9:	9.		2	8	. 2
SabSTAR(Y92A)	91	120	2	ຄ	₹	=	2	_ £1	7.3 18	₹ -	1.7		2	-	0.0	0.5	_	š	8	*
SakSTAR(E91A,K94A)		6	18.2	2	=	9.	~~	€		۰	_		: =					:	:	•
SakSTAR(K94A,N95A,K97A)	72		ž													•		š	3	5
SubSTAR(N95A)	n	992	2	=	4.0	2	=	8	13	÷		- 2	2	£,	5.9		***	: 2	: 2.	. g
SakSTAR(K96A.K97A.K98A)		÷	12.8		2	37	-^	×16	9.1	2	2	0.58		2			0.30			
SukSTAR(E99A)	2	- 7	~	5	0.4	7	8.9	. 22	2.7 4.7	٠ <u>۾</u>	<u>8</u>	-7	6.2				80	6	5	2
SakSTAR(E99A.E)00A)	11								•					!				!		
Sakstaritiona	ន	£	9.	=	7	9	2.	2	2.8 7.6		9		\$.	2.4	67		8.0	. 20	8	\$
-	-		_				_					_	,	1			-	:	ł	

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	-				$\ \ $					munine MAbs	MABs	$\ \ $					4			1
Variation	EP.	Spec. Act.	1157/1	26.6.2 10.6.2 28.12 10.10	cluster 0.K7	12		HET 71481		cluster !!	2		23E	ed Po	LOCS 24C4	2	900	SakSTAR pedent plasma Subnool B Subn	n plasms Subroof C	1
Sakstar(K102A)		í fig		=	1 23	2		F			ف	Ē.	-	Ê	e	86		ió	2	1
SukSTAR(S10)A)	-50	210	0.6		5.0 9.4	1.9	<u>e</u>	8.9	=	2	3.6	<u>8:</u>		\$	78	6:0	3	8	S	•
534STAR(F104A)	<u> </u>	ž	8.8	•	1.8	1 27	2.2	\$0	7	4.	ਵ	3	1.6	2	=	2	\$	8	2	
Salstariiloa)	<u></u>	5	ដ	· •	3.0 7	7.4 6.7	5.5	3	=	=	2	8 9	Ξ.	8 9.	7	S	8	8	2	
SetSTAR(T107A)	77	8	5.2	15	3.4 9.8	9	~	8.7	4.7	I	6:	-5	63	3.2	9.0	8.0	z	z	86	
SakSTAR(E108A.K109A)		2	911	5.1	7.2 19	2	*	=	7	=	7	0.43	6.9	3	2	6:				
SukSTARIFI11A)	~	\$	3.7	91	3.8 13	2	5	2	=	3.1	9.0	5.8	2.9	.	2	6.0	8	8	86	
Salstarivilla.VIII)AI	3	951	₽		3.9 10	~	Z.	5.8	=	8.0	0.3	<u> </u>	3	2.3	3.0	8.0	98	2	95	
SakSTAR(D) 15A.S117A)	98	×	33	=	5	2	-	3.	•	63	6	<u>.</u>	8,4	2.6	2	6.0	8	2	8	
SutSTAR(D115A.E118A.H119A)		32	<u> </u>	32 3.4	7	8.7	=	6.6	8		1.2	<u>.</u>	*	7.7	0.6	8.				
SakSTAR(L116A.S117A)	2	\$		35 3.6	6 33	7	8	8	220	ę	é	0.3	Ţ	6;	ž	9:	š	2	86	
SahSTAR(H119A,K121A)		85	18.0	=	2 -	8	3	ž	2	2	0.52	2	=	2	2	71				61
SzkSTAR(II 20A)	92	75	2	36	3.1	9	2	8.6	ສ	0.6	6.9	3.0	2	2.2	23	0.	2	8	88	
SakSTAR(N123A)	<u>.</u>	•	¥														8	8	\$	
SukSTARIF125A)		410	2.8	13 4.7	-	=	=	22	6.0	6.	å.	3	23	7	6.9	9.1	8	8	8	
SakSTAR(N126V)	<u> </u>	15	1.6	13 . 20	2 0	=	2	85	8	8.8	ສ	• •	0.0	Ş	6.5	0.7	86	8	88	
Sakstaril17A)	Ξ.	75	8.0	£1 E8	8 5.0	9.9	×	\$	=	2	2	6.0	<u>e:</u>	60	Ħ	=	5	z	86	
SakSTAR(1128A)	•	<u>-</u> ۾	16 2	25 4.8	æ 2	=	ĕ	7.6	\$	8.2	5.9	ຊ	2	3	6.7	6.0	8	8	8	
SukSTAR(T129A)	4	8	1 (3	2.3	÷	*	<u>~</u>	=	2	7	1,1	0.7	2	2	5	0	8	.8	\$	
Sakstar(K130A)	130	280	1.	3.2	4.0	3.5	2	6.7	=	2	1.1	ğ	ą	7	6.0	9.0	6	2	7	
SakSTAR(VI31A)	130	٥	6.3	1 2.9	=	=	<u> </u>	Ξ	2	2	Z	<u> </u>	2	3	9.8	6.0	8	8	25	
SetSTAR(V132A)	8	130	7	3 2.6	5 9.2	=	3	2	8	2	17	7.1	3.6		3.6	9.0	8	8		
SakSTAR(II)3JA)		8	7.6	\$ 1.9	9.7	8. 7.8	₹.	6.0	-	9:	<u> </u>	0.56	7.	9.1	9.1	6.0	8	2	66	
S#STAR(E134A.K135A.K136A)		:	2 22	1 6.7	22	22	*	2 2	2.	27	1.7	0.2	=	3.0	9.0	2.61				
SASTARIKINSA	\$4	410	3.2	2 13	7.9	=	8	=	=	3.8	2.0	9.	6.9	3.7	<u>°:</u>	6.0	8	2.	šė	
SakSTARIKINAA							-										•			

Table 4: Mutagenesis of S34, G36 and H43: Association constants (K, x 10'mol/L') for binding to insolubilized murine monoclonal antibodies (Mab) SakSTAR patient plasma 0.5 ₿ å é. é é and absorption (percent) of antibodies of immunized patient plasma 17G11 26A2 30A2 2B12 3G10 7 -Spec. Act. (RU/mg) 2 (mg/L) Ę. ek\$TAR(\$;40,036R,H43R,K74A) #STAR(\$34G,@36R,H43R) Sat STARIG36R.K74A.K135R) SakSTARIG36R.K74R.K135R) ALSTARIG36R.K74A.N9SA) akSTAR(S34G,G36R,K74A) akSTAR(K35G,G36R,H43D) akSTARIG36R.K74A) LASTAR(Q36R,K74R) #STARIS34G.G36R) SASTAR(S.WA) JASTARIG16A) ASTARIG36E ak STARIG36K) JASTARIG36N) LLSTAR(H43R) ANSTARIG36L) akSTARIG36Q) ASTAR(036R) ASTAR(H43A) Variant

Table 5: Mutagenesis of K35, Y73, K74, E80/D82, N95, K130, V132 and K135: Association constants (K, x 10'mol/L.') for binding to insolubilized murine monoclonal antibodies (Mab) and absorption (percent) of antibodies of immunized patient plasma Subpool 2 2 3 2 8 93 2 2 2 8 2 z 2 SakSTAR patient plasma Subpool B 2 S 2 2 2 5 2 2 8 z 3 2 8 8 2 40C8 24C4 1A10 9 = 2 2 6.0 60 9 S 2 7 6.0 9.13 é 章 ŝ 6. 2 ę ₹ 50 9 6 Ç Ħ ᅙ 8.9 를 ₹ ŧ ę ę é 5 20 BF12 14H5 28H4 32B2 7F10 2 5.6 7 20 7 ~ Ξ 2 2.7 2 3 ឧ 6 2 3. <u>6</u> é 夸 JOAZ ZBIZ 3GIO 0 = # Ξ 5 7 : 7 5. 3.6 ē 를 ę. **8** 6. 2 5 3 2 2 7 3 . 9.0 Epitope cluster é 章 를 2 Ξ 0,7 2 2 3.2 17GH 26AZ 6 = 2 9.0 2 2 Spec. Act. (RU/mg) 2 8 2 5 ŋ 8 3 8 3 ŋ v ٥ ٥ 2 8 8 (mg/L) Ë AKSTAR(S340,G36R,H43R) ekSTAR(E80A,D82A) SakSTARIY73Wı SalsTAR(K35B) SakSTAR(K35Q) SALSTARIY7,1F1 SakSTAR(Y73H) SakSTAR(Y73L) SakSTAR(K74N) SakSTAR(K74Q) SakSTAR(K74R) SukSTAR(Y735) SakSTAR(K74E) SakSTAR(EBOA) SalsTAR(D82A) SakSTAR(N950) akSTAR(K3SA) SakSTAR(N95E) MSTARIY73A) akSTAR(K74A) SakSTAR(N95K) S.ASTAR(N95R) SETARIN95A) Variant

,	•	,							manne	500		l							
HT.F.	Erp.			Epito	Epitope clusier		\vdash	ľ	Epriope cluster II	aster II	Γ		Epilo	Epitope cluster III			Š	SakSTAR puttent plasma	d plasma
	(mg/L)	.) (kWmg) 17GT	17611	76A7	200	30A2 2812 3G10 18F12 14H5 28H4 32B2		11 TH	2334)F10		1361	E	40CB 24C4 1A16	OLX.	1000	Subpool B	Subpool
Saksta R(K130A)		780		13	3.2 6.	6.4 3.5	~	.9	=	2	72		é	;	:	*	8	7	
SakSTAR(K130T)		230		2	3.5	80 5.6	<u> </u>	3.	9.7	: =		é	9				: 6	: 2	: :
Salstar(V132A)	2 2	81	<u>.</u>	5	2.6 9.2	= ~	_=	2	8	2			3.6	_		*	: 2	. 2	: ;
SekSTAR(V132L)	136	120	9	-	2.3 8.0	7.6 0	63	2	4	2	: :	2.0	=	6	60		: 5		8
SatSTAR(V13ZT)	82	3	2	2	24 7.8	0.6	<u>8</u>	2	ສ	2	_= ?	=	20		_			: ;	: ;
SekSTAR(VI32N)	_9	051	÷.	=	1.7 7.0	2,7	_=	-	2				: 2			-		: ;	2 8
SaLSTAR(V132R)	<u></u> %	82	*	~	08	32	_ =	\$3	7.8								: 8	: :	2 8
SakSTAR(K13SA)		410	5.2	=	9,7	=	2	=	=	80						- 8	: *	: 8	: 8
SakSTAR(K!35F)	8	3	3.9	6.3	- -	7	_=	Ĵ	2	<u> </u>						-	: :	: 8	
SakSTAR(K1.15R)	<u> </u>	238	0.4	- - -	6.0	50	_=	=	=	2	<u>~</u>	7.	7.	-				: :	: 8
SakSTARIKJSA.K74A)	유	130	Ē				_		:							<u> </u>	2 2	:	: :
SakSTAR(Y73A.K74A)	_ *	ŋ	=	.e.	 	6	2	63	2	9	<u></u>	_		-	-				
SakSTAR(Y7.)F.K74A)	<u>.</u> 5	•	2	5. 6.	77		==	2	=				_	•			; ;	; 3	. 8
Sakstarii60a.K74a.N95ai		:	2	2.7 cb.1	2.5	53	=	11	9.6	3.4	<u></u>				6.0		: 8	: \$. 2
SakSTARIN9SA.K135R)	130	740	o.	6.1	9.0	6.6	_=	=	2	2	_ <u>=</u> _	*					· ×	8	. 26
SALSTARCK! 30T.K! 35R)	<u>s</u>	280	3.7	9.1	7.7	80	=		Ž,	3.7	9		, . E				. 2	: 5	
_								:	•							_	À	8	

Table 6: Combination mutants of SakSTAR(K130T,K135R) with K35A, G36R, E65X,K74X and selected other amino acids

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SY41 SY41 SY3

SY71

5Y59 5Y51 5Y49 SY 35

SY6S SY66 SY67

SY 36

5Y57 5Y70 5Y58

c0.1 2.4

325TAR(E65Q.K74Q.T90A,K130A,K133R)

ASTAR(E65Q.K74Q.K130A.K133A) ASTAR(K74Q.K130Q.V132R.K135R)

ALSTAR(E65Q.K74Q.K130A.K135R)

42TAR(K74Q,K130E,K135R)

. 19

SY 69

Subpool C Pool 40 SakSTAR patient plaxm 25E1 40C8 24C4 TATO **6.1** 3.5 9.1 ę é TAHS ZBH4 3ZBZ Erp. 546STAR(K74Q.K130T.K135R.K136A.+137A) 445TAR(Q36R,H43R,K74R,K130T,K135R) SakSTARIGNGR.E65A.K74Q.K130T.K135R1 5.4STAR(534G,G36R,K74Q,K1,30T,K1,35R) SakSTAR(G)6R.E6SA.K74A.K1,10A.K1,15R1 SokSTAR(E65A.A72S.K74Q.K130T.K135R1 atSTAR(E65Q,K74Q,E75A,K130T,K135R) iakSTARIE68Q.K74Q.E75D.K130T.K135R1 ALSTAR(E65Q.T71S.K74Q.K130T.K135R) MASTARIG36R,K74R,K130T,K135R) ASTAR(G36R,K74Q,K130T,K135R) 14KSTAR(E6SA,K74Q,K130T,K135R) 45TAR(K74Q.K86A.K130T.K135R) 5at STAR(E65Q.K74Q.K130T.K135R) ASTAR(G36R,K130T,K135R) BASTAR(K74R,K130T,K135R) ASTAR(K74Q,K130T,K135R) 45TAR(K74Q,K130A,K135R) AESTARKKI SØT, KISSR)

SUBSTITUTE SHEET (RULE 26)

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Variable		Spec. Act		Forto	Fostone chatter		-	mutine with the chaire I Enline classes I Eo	attoo of			1	I\$	Epitope cluster	F	\dagger	TES.	STAR palle	M plasms	
	3	(kU/mg)	1101	78X3	ZOXZ	E ZIE	<u> </u>	F17 14	F 28	1	94	E	i		Ę	er. K	20110 Sal	S Bloods	40C8 24C4 1A10 Pool 10 Subpool B Subpool C Pool 40	9 1 1 1 1
SALSTAR(E6SQ,K74Q,N9SA,K130A,K135R)	ę	220	Ļ	-	2	Ē	F	Ė	*	þ	þ	Ę	Ę	E	F	F	E	ê	F	34.12
SatSTAR(E65Q.K74Q,E11tA,K130A,K135R)	8	2	8.5	<u>=</u>	2.8	15 27	=	7	5.7	2	2.6	 9	é.	6.1	2.8	6.5	8	22	r.	S8 SY73
Sabstar(E61Q.K74Q,N95A,B118A,K130A,K135R)	ä	8	7.8	=	7.7	12 21	2	3.9	6.1	9.6	ລ	-	é	5.8 2	2.5	<u>ა</u>	\$	11	74	S8 SY74
SakSTAR(NOSA,K130A,K135R)	2	9	9.1	=	.	=	3.	5.9	9.6	6.8	22	6	E	4.5	3.0	9.0	8	=	82	<u> </u>
S#STARIK13A,E63Q,K74Q,K130A,K133R)	2	2	Ę														ŧ	76	3	45 SY75
S=ASTAR(K35A,H43R,E63Q,K74Q,K130A,K133R)			ž														\$	22	22	SS SY76
SakSTARIE65Q.K74Q.S103A.K1,30A.K1,33R1	Ħ	8	6.7	2	7.6	2 19	8	7.7	3.9	6.3	ລ	-	ē.	1.6	0 9.1	9.0	55	n	2	61 SY77
Sanstar(T21A.K.)SA.E65Q.K74Q.K1,10A.K1,15R1		•	Ę				-										8.	36	22	50 SY78
\$4\$TAR(T36A,E63Q,K74Q,K130T,K135R)		981	¥														5	<u>.</u>	19	55 SY79
SakSTARIK37A.ES8A.E61A.K74Q.K1.30T.K135R)		120	¥														5	*	5	54 SY80
S44STAR(E65Q,K74Q,K109A,K130T,K135R)	Ş	310	٤,٢	25	7.1	12 12	_=	22	0,	5.8	ລ	5	<u>é</u>	3.4	8: 0		2	=	89	51 SY81
SakSTAR(E65Q,K74Q,E108A,K1,0T,K1,15R)		92															5	*	2	SY 82
SaLSTARIE65Q,K74Q,E108A,K109A,K130T,K135R)	7	2	5	=	_	13 17	-	3.0	7	8.8	ສ	를 -		3.7 2.	2.6 0.5	ده.	83	12	19	50 SYB3
SabSTARIE65Q.K74Q.K121A.K130T.K135R)	2	150	5.7	=	5	=	2	1.	4.6	.	ē	ě		3.5	1.8 0.9		5	22	69	57 SYBS
SabSTARIEI9A.E65Q.K74Q.K130T.K135R)			¥														15	11	70	SY B6
SabSTARIE65Q.K74Q.D115A.K130T.K135R1		23	Ę														25	22	63	SY87
SAASTARIG10R.E65A.K74Q.KI,10E.VIJZR,KI,15R)	÷	8	3,6	-	_	=	-	2	1	\$	0	9.			46.1 0.9	ο.	3	1	5	44 SY60
SA\$TAR(E65Q,K74Q,N95A,E118A,K130A,K135R,+137A)		2															\$	30	2	60 SY93
Sabstar(E63Q.K74QJN95A.E118A.K130A.K133R.K136A.+137K)		1,400									1						3	2	2	75 Z

Association constants ≥ 10-fold lower and antibody absorption ≤60 percent of wild-type SakSTAR are represented in bold type; ≥ 100,000 HU/mg represented in bold type. NT: not tested.

Table 7: Combination mutants of §	SakST	FARU	380	A,D8	12A,	K130	I,K	135R) wit	ξ.	5A, (361	Ä	5X, F	74X	, and	l selecí	ed othe	SakSTAR(E80A,D82A,K130T,K135R) with K35A, G36R, E65X, K74X, and selected other amino acids	acids	
			L					П	Jan.	murine MAbs		$\ \cdot\ $	ľ	Taises chistory	111			SakSTAR patient plasma	tieni plasma	Γ	
Variant	Exp. (mg/mt.	Spec. Act. (kU/mg)		17G11 26A2	A7 30A7 2B	JOAN 2BIN	2,3010	TEEF		TAKS ZEHA 3ZBZ	Ĺ.	TETO THI	Ľ	23E1 40C8		¥.	Pool 10	Subpool B	Subpool C	Pool 40	8
	-	-		F	ŧ	F	-	_	Ļ	1		5	1	Table 1	Ę	F	2	3	20		286
Sakstangadazazakijot, kijsk)	₹.	3 5		: =		: =	: =	2	2	32	3.9	₩.	6	- ¢	6	%	7	×	8	ני	SYJ
SASTAR(K74R,E80A,D82A,K130T,K135R)		P	3;	. :		2 5	: ;	: :					4	<u>6</u>	Ą	8.0	4	2	8	4	SYIS
Sakstar(K74Q,E80A,D82A,K130T,K135R)		<u> </u>		6	2 :	2	3 :	1 1	; :	_				· -	4	8.0	8	*	8	28	SYI7
Salstarik.35a.K74R.E80a.D82a.K1.30T.K1.35R1	2	<u>s</u>	<u>ç.</u>	5.	č	2	ž	2	2					•	5		7	=	3	25	SY 19
SakSTAR(E65D.K74R,E80A,D82A,K130T,K135R)	¤	2	3	2	5.9	\$	=	<u> </u>	=	~ ~	- -	ਵ ਵ		3	3	3	; ;	: :	: \$		SY 20
SakSTAR(E655.K74R.E80A,D82A,K130T.K138)		91	2	≃	9.	1	•	2	2.0	5 0	1 9.9	₹ <u>-</u>	<u>é</u>	ē -	Ę	7 .	S	2	8		
Sabstabless K248 E80A D82A K130T K135R)	2	7		6	9.9	3.6	g	22	2	8.2		2.0 db.1	<u>8</u>	-0.1	ᅙ	<u>o</u>	86	*	9		24.21
COLLY THE WASHING CLEAR THE AND THE WASHINGTON		350		Z	2.9	2	×	2	8.9	18 5	5.5	0.1 6.1	<u>.</u>	8	6	60	22	æ	5 5		SY:0
-	ç	9	<u> </u>	9	=	5.7	\$5	_ ~	2	=	2	2.2	<u>8</u>	9.	6.	1.0	5	11	8		SX I
	? :				=	20	*	\$	2	. ~	2 6.7	- 7.	- 48.	 6	ફ	6.0	\$	2	69	•	SY23
	: :		9		· •	0.0	8	_ ∞	2	-	2	23 AB	£.	4	Ą	8.0	3	=	\$	8	SY22
	; ;						=	9	7.8	9	-	- 6	.a.	18	Ą	0.7	22	3	G	2	SYE
	; ;	<u> </u>	:	: :	: :			9	=		36	3. A	£.	1.60.1	章	1.2	5	2	8	Z	SYS
(36)	× - 2	7					: :	. 2	: #		6.8	22	 	ş	ŧ	6.0	2	2	2	#	8730
Sec. 1 A (E02D, R /4Q, E00A, D8.1A, R 30 R	3 3	2 5	3 7		3 2		: 2	2	. 8	24		 &	.e	6	6	0.1	\$	=	ž	7	SY47
135R)	; <u>\$</u>	2	<u> </u>		2,	•	2	2	-	=	7	2. 60.1	<u></u>	.	Ą	0.1	38	-	8	\$	SY 46

Table 7 - cont'd: Combination mutants of SakSTAR(E80A,D82A,K130T,K135R) with K35A, G36R, E65X, K74X, and selected other amino acids

			۱	ı					munne N	N S			l	l		Γ					
Verlant	Esp.	Spec.		4	Epitope cluster	īg.	Γ		Epitope cluster II	luster II		-	Ħ	Epitope cluster III	rier III		L	SakSTAR p	SakSTAR patient plasma	_	
	(mg/mL)	(mg/mL) (kU/mg) 17011 26A2 30A2 2B12 3G10 18F12	100	76A7	30.42	71817	3010	TBFTZ	14H5 28H4 32B2 7F10 7H11 25E1 40C8 24C4 1A10	1				2	Ĕ	אַנ	700 TO	Subpool B	Pool 10 Subpool B Subpool C Pool 40	P00140	ð
SAESTARIR71R-EBOX DBZAS1035.K139T.K135R)	E	89	E	F	E	Ŀ	6	2	17 63	E	F	T	त्वा त्वा	¢∰.	Ę.	6	6	ř	66	þ	243
S485TAR(K15A,E65D,K74R,E80A,D82A,E108A,K109A,K130T,K135R)	. 0.6	ŝ	5.8	2	2.6	92	2	=	16 13	3.2	=	ŧ	€	<u>6</u>	Ą	0.5	88	2	8	ŧ	SYI2
SakSTARIK;1SA,E65D,K74R,E80A,D82A,E108A,K130T,K135R)	2	5	¥.	2	3.0	≃	3.9	22	17 7.4	7.	6.	₹	₹	4	章	6.0	3	•	8	8	SY32
SASTARIESSO, KTIR. E80A, D82A, E108A, K110T, K1,15R1	-	8	Ξ.	6.7	6.9		8	6		=	7.1	₹	ફ	章	ģ	0.1	8	=	8	•	SY33
SakSTAR(K?3A.E65D.K?4R.EB0A.D82A.K190A.K130T,K133R)	7	2	\$3	=	53	=		=	17 21	2	[]	<u>6</u>	é	Ē	6	<u>e</u>	\$	v	5	8	SY36
S&STAR(E65D.K74R,E80A,D82A,K109A,K130T,K133R)	8	8	9.7	9.6	8.9	7	28	=	32 12	=	2.3	é	6	é	Ę	6.0	*	2	3	23	SY37
S#ETAR(K15A_E65D.K74R.E80A,D82A,K130T,K135R,K136A)	88	=	:	~	33	=	- []	22	13 7.6	\$	9.	é	Ę	é	ŧ	8.0	\$	=	23	\$	SYM
Sabstar(E65D.K74R,E80A,D82A,K)30T,K)33R,K136A)	\$	<u> </u>	8.9	5.8	3	\$.	2	33	32 14	7.9	2.0	é	é	6	é	9.8	*	22	69	.\$	SY35
SakSTAR(E65Q,K74Q,D82A,S84A,K130T,K135R)		2	ž														\$	17	\$	\$	SYSON
\$48TAR(K35A.E65D,K74R.E80A.D82A.K86A.K130T.K133T)	89	<u>`</u>	;	2	5.5	51		2	12 6.4	6.3	6.	₹	é	9	ө	0.	*	•	\$	\$\$	SY40
S4LSTAR(K33A,K74Q,E80A,D82A,K130T,K133R)	2	2	6.1	ž	23	3.0	5.9	38	9.8	8.9	6:	₹	é	₹	<u>é</u>	9.0	\$	2	3	#	SYZB
8415TAR(K35A_E61D,K74R_E80A,D82A,K130T,K135R)	z.	2	_	2.5	6.9	5.5	ม	37	7.	1.7	23	ę	ą	ę	章	0.1	3	82	5	\$\$	SY 29
SakSTAR(K35A,E65D,K74R,E80A,D82A,V132R,K135R)	=	×	6.7	23	5.3	2	2		61 61	2.	2.0	6	ą	章	é	=	3	2	80	3	SY61
\$4\$\$TAR(K35A.E65D.K74R.E80A.D82A.T129A.K135R)	=		7.0	=	3.	=	~	23	= =	6.3	2.2	₹	鱼	ę	륟	6.	×	=	2	8	SY62
SakSTAR(K35A,E65D,K74R,E80A,D82A,T129A,K135A)	្ន	 	6.9	#	3.8	22	R	39	6.6 9.7	×	2.1	इ	é	春	章	6.0	*	=	5	8	2764
	_	_					_					_				_	_		:		

Association constants ≥ 10-fold lower and antibody absorption ≤60 percent of wild-type SakSTAR are represented in bold type; ≥ 100,000 HU/mg represented in bold type. NT: not tested.

Table 8: SakSTAR variants with intact specific activity (> 100 kHU/mg) and 550 percent absorption of human antibodies elicited by treatment with

Variant	Spec. Act.		SakSTAR p	SakSTAR patient plasma		
v ditait	(kU/mg)	Pool 10	Subpool B	Subpool C	Pool 40	Code
SakSTAR(K74Q,K130T,K135R)	130	20	25	19	62	SY41
SakSTAR(E65A,K74Q,K130T,K135R)	170	45	16	11	22	SY48
SakSTAR(E65Q,T71S,K74Q,K130T,K135R)	210	49	21	2	29	SY65
SakSTAR(E65Q,K74Q,E118A,K130A,K135R)	180	20	78	72	28	SY73
SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R)	190	48	7.7	74	28	SY74
SakSTAR(K35A,E65Q,K74Q,K130A,K135R)	110	6	56	63	45	SY75
SakSTAR(E65Q.K74Q,K109A,K130T,K135R)	210	20	22	89	51	SY81
SakSTAR(K74Q.E80A,D82A,K130T,K135R)	110	46	17	09	8	SYIS
SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R)	140	43	11	89	57	SY19
SakSTAR(E65S,K74R,E80A,D82A,K130T,K135R)	110	35	12	09	•	SY20
SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R,K136A)	100	46	78	<i>L</i> 9	45	SY35
SakSTAR(K35A,K74Q.E80A,D82A,K130T,K135R)	120	6	16	2	48	SY28
SakSTAR(E65D,K74Q.E80A,D82A,K130T,K135R)	110	43	13	2	42	SY30
SakSTAR(E65Q,K74Q,E80A,D82A,K130T,K135R)	120	43	21	2	42	SY47
SakSTAR(E65Q,K74Q,D82A,S84A,K130T,K135R)	170	45	21	09	45	SYSON
SakSTAR(K35A.E65D.K74Q.E80A.D82A.K130T,K135R)	140	35	œ	58	40	SY46
SakSTAR(T21A,K35A,E65Q,K74Q,K130A,K135R)	110	20	7 6	72	20	SY78
SakSTAR(E65Q,K74Q,K109A,K121A,K130A,K135R)	140	20	31	73	52	SY88
SakSTAR(F650 K740 D824 S844 K1004 K1304 K135R)	180	43	20	62	4	SY89

Table 8 - cont'd: SakSTAR variants with intact specific activity (2 100 kHU/mg) and <50 percent absorption of human antibodies elicited by treatment with wild-type SakSTAR

יו כשוווכוור אוווו אווס-וללי-סשט זיטיו						
Variant	Spec. Act.		SakSTAR patient plasma	ient plasma		
	(kU/mg)	Pool 10	(kU/mg) Pool 10 Subpool B Subpool Pool 40	Subpool C	Pool 40	Code
SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R,V137A)	120	120 45	30	74	60 SY93	\$793
SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R,K136A,V137K) 1,400	1,400	37	16	70	54	SY94
SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130A,K135R)	110	46	76	63	41	SY95
Antibody absorption <60 percent of wild-two SakSTAB are represented in hold two: > 100 000 HIV/ms represented in hold type	i bod d	و. ×	000 H 1/m	representer	t bold to	ģ

Table 2: Fibrinolytic properties of selected SakSTAR variants in human plasma in vitro

<u>(</u>(1)

Compound	Fibrinolytic potency (C50 in µg/mL)	Residual fibrinogen at C50 (% of baseline)	Fibrinogenolytic potency (C50 in µg/mL)	Code	
SakSTAR	0.18±0.01	93±3.5	24 ± 3.6		
SakSTAR(K74Q,E80A,D82A,K130T,K135R)	0.15 ± 0.01	97 ± 3.0	14 ± 3.2	SY15	7
SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R)	0.24 ± 0.04	94 ± 10	29 ± 3.1	SY19	71
SakSTAR(K35A,E65D,K74Q,E80A,D82A,K130T,K135R)	0.11 ± 0.01	92±3.0	20±2.0	SY46	
SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R,K136A,V137K)	0.13	16		SY93	
W. SR					

The data represent mean ± SD of 3 experiments. C₅₀: amount of wild type or variant SakSTAR required for 50% clot lysis or 50% fibrinogen breakdown in 2 hrs.

<u>Table 10:</u> Pharmacokinetic parameters of the disposition of staphylokinase-related antigen from plasma following bolus injection of SakSTAR

Variant C ₀ (µg/mL	C ₀ A B (1/2 (α) (1/2 (β) ($\mu g/mL$) ($\mu g/mL$) (min) (min)	В (µg/mL)	t1/2 (α) (min)	t1/2 (β) (min)	VC (mL)	AUC Cip (ug.min.mL-1) (mL.min-1)	Cip (mL.min ⁻¹)
SakSTAR 0.8±0.	0.8 ± 0.1 0.6 ± 0.1 0.2 ± 0.0 2.8	0.2 ± 0.0	2.8	7.0		1	2.2 ± 0.2
SakSTAR(K74Q,E80A,D82A,K130T,K135R) 0.5±0.	0.5±0.1 0.4±0.1 0.1±0.0	0.1 ± 0.0	2.0	9	20±2.2	2.5 ± 0.3	4.1 ± 0.5
SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R) 0.6±0.0	0.6±0.0 0.5±0.0	0.1 ± 0.0	2.0	2	16±1.1	2.8 ± 0.2	3.7 ± 0.3
SakSTAR(K35A,E65DK74Q,E80A,D82A,K130T,K135R) 1.1±0.1 1.0±0.1 0.1±0.0	1.0 ± 0.1	0.1 ± 0.0	2.0	24	9.6 ± 0.7	6.4 ± 0.5	1.6 ± 0.1

Additional therapy			Stenting Ich IP artery	Right upper leg amputation	PTA	ojmosoutis tedem 1	Cumoal sympathic connic	Desobstruction	¥1.4	Left AF graff	PTA	•	•	. !	New right FP graft	•	PTA + stenting	FF graft			İ	X1	Summer	A TOTAL CONTRACTOR AND	Aspiration infomoccomy, FTA	5552	1_			Pseudo aneurysm, right AF graft	revision	•		• • • • • • • • • • • • • • • • • • • •	Aspiration thrombectomy	¥L.	15±4.3 19±9.4 12±2.8 14±4.4
Total duration of infusion		;	57	23	7	20	2 :	2 5	2 •		n •	n ;	ę .	n ?	4 :	£ :	<u>.</u>	\$	19±3.5		7	: .	ş	,	, ×	=	20 ± 4.0			•	ş	7 -	₹ -	4,	; ه	-	14±4.4
Total dose of thrombolytic agent (mg)	8	•	7 :	=	00	22	2	۰ (, , ,	3 -	* 4	9 9	<u>.</u>	-	3 2	3:	2 5		13 ± 2.1		. 90	<u>«</u>	2	<u>-</u>	<u> </u>	8.5	16±3.4		,	80	4	2 2	2 4	۰ ۵	× 7	67	12 ± 2.8
Recanslization by thrombolysis		of the second	מוווחובוב	Partia	Complete	Partial	Complete	Complete	Complete	Complete	Partial	Complete	Complete	Complete	Complete	Complete	Complete	Complete			Complete	Complete	Complete	Partial	Complete	Complete				Complete	Complete	Complete	Complete	Compress	Complete	Complete	
Length of occlusion (cm)		•	•	۰ ۰	n	S	<u></u>	2	50	•	· •c	, <u>s</u>	: -	. *	1 S	3 5	2 5		18 1 3.5		•	œ	6	*	ø	15	18 ± 10		•	×o	9	<u> </u>	2 •	> 5	2 ∝	٥	19 ± 9.4
Age of occlusion (days)		×		ų <u>s</u>	2 ;	8	_	~	4	7	_	4	4		-		· oc	16444	0.0 I 2.1		90	7	Š	4	2	2	13±4.3		,	ŧ	7	7	21	; ×	3 8		15±4.3
Locus of Age of Length of Recanalization Total dose of Total occlusion occlusion by thrombolytic of (days) (cm) thrombolysis secun (mp)		Femoro-femoral graft	Riohr PA	A 20 CEA	איין יויין	Kight SFA	Right AF graft	Len FT gran	Right IF graft	Left AFS	Left tibial anery	Right FP graft	Left radial anery	Right FP graft	Left PT graft	Len FT gran	Right SFA graft	,			Len SFA	Right C.I.A.	Right E.I.A.	Left FP junction	Len SFA	Right FP bypass		á	A) Picht E ! A	, C. 1. 2. 1	Right AF graft	Left anterior tibial anery	SFA	Left PA	Right SFA	١.	•
Clinical		Subacute	Acute	Rethain	Tipodica.	Supacure	Claudication	Subacute	Restpain	Acute	Restpain	Claudication	Acute	Acute	Acute	Claudication	Restpain			30T,K135R)	Claudication	Subacute	Acute	Claudication	Restpain	Acute		SakSTAR(E65D,K74R,E80A,D82A,K130T,K136B)	Subacute		Restpain	Restpain	Restpain	Subacute	Claudication		
Age (vrs)		99	23	63	4	? :	ř	2	8	æ :	67	\$	~	2	57	8	11	62 ± 3.1		D82A,KI.	છ :	;	7 5	2 (3 8	٩	39 ± 4.7	80 A. D.82.	57		8	2;	9:	6 5	92	67±3.4	
Gender		Σ	Σ	>	Σ	u	L 6	٠;	Σ	> ;	Σ	Σ	Σ	æ ;	Σ	Σ		· ~		Q,E80A.	Σ	Ξ >	ΞΞ	Ξ 2	Σ 3		5	D.K74R.E	Σ		Σ:	٤٤	<u>ب</u> د	٠.	Σ	I Si	i
Patient 1d.	SHOLAK	15	VERM	GEI<	Ž	8	3 =		ביים ביים ביים	Š	Z Z	AXIV	HZY.	*	SKA SKA	Z C	CAM	Mean # SEM		SakSTAR(K74Q,E80A,D82A,K130T,K135R)	47×	<u> 2</u>	STRO	VEDC	S H	Mean + CEL	I Call	*KSTAR(E6S)	URB		NO.	7 20	Y	₹ • č	-	Mean ± SEM	

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with SakSTAR variants in patients with peripheral arterial occlusion

Treatment Absorbant	SakSTAR		Insolubilized compound SakSTAR(K74Q,E80A,D82A,K130T,K135R) SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R)
SakSTAR (Pool 40) SakSTAR SakSTAR(K74Q,E80A,D82A,K130T,K135R) SalSTAR(E65D,K74R,E80A,D82A,K130T,K13SR)	95 48 57		
SakSTAR(K74Q,E80A,D82A,K130T,K135R) (Imb Vin., Ver Gie.)	r Gie.)		
SakSTAR SakSTAR(K74Q,E80A,D82A,K130T,K13SR) SalSTAR(E65D,K74R,E80A,D82A,K130T,K13SR)	94 91 92	95 93 94	95 89 94
SakSTAR(E65D,K74R,E80A,D82A,K130T,K135R) (Urb.)			
SakSTAR SakSTAR(K74Q,E80A,D82A,K130T,K135R) SalSTAR(E65D,K74R,E80A,D82A,K130T,K135R)	8 2 2	88 93 85	85 94 94
Data represent median values of the percent absorption with 250 nM absorbant, measured by residual binding to insolubilized compound.	0 nM absorban	t, measured by residual binding to insolubilized coi	mpound.

Table 13: Additive substitution mutagenesis of SakSTAR(E65Q,K74Q,K130T, K135R) with selected other amino acids

Variant	Spec. Act.	Antibody	Code	
	(kU/mg)	absorption (percent)		
akSTAR(E650,K740,K130T,K135R)	150	\$9	SY49	
SakSTAR(E65Q,K74Q,D82A,S84A,K130T,K135R)	170	45	SY 50	
SakSTAR(E65Q.K74Q.T90A.E99D.T101S.K130A.K135R)	410	518	SY98	
SakSTAR(E65Q.K74Q.E108A,K109A,K130T,K135R)	180		SY83	
SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130T,K135R)	110	41	SY95	
SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130T,K135R,K136A,V137K)	1,500	90	SY118	75
SakSTAR(E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,K135R,K136A,V137K)	2,900	78	SY128	
SakSTAR(K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,K135R,K136A,V137K)	3,700	24	SY 141	
SakSTAR(K35A,E65Q,K74R,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K130T,K135R,K136A,V137K)	5,700	31	SY145	
	-			

Spec. Act. 2 100 kU/mg is represented in bold type. Absorption of antibodies (in percent) from pooled immunized patient plasma; values 560% are represented in bold type.

Table 14: Fibrinolytic properties of SakSTAR variants in human plasma in vitro

Compound	Fibrinolytic potency (C50 in µg/ml)	Residual fibrinogen at C50 (% of baseline)	Fibrinogenolytic potency (C50 in µg/ml)	Code
SakSTAR	0.18 ± 0.01	93±3.5	24±3.6	
SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130T,K135R,K136A,V137K)	0.15 ± 0.02	90 ± 5.0	14 ± 1.0	SY118
SakSTAR(K3SA.E6SQ.K74Q.D82A.S84A:T90A.E99D.T101S.E108A.K109A.K130T.K135R.K136A.Q137K)	0.17 ± 0.01	87±3.0	7 ± 0.6	SY141
SakSTAR(K3SA.E6SQ.K74R.D82A.S84A;T90A.E99D.T101S.E108A.K109A.K130T.K135R.K136A.V137K)	0.19 ± 0.01	82 ± 3.0	7 ± 0.9	SY145

The data represent mean ± SD of 3 experiments. C50: amount of wild type or variant SakSTAR required for 50% clot lysis or 50% fibrinogen breakdown in the absence of fibrin in 2 hrs.

Patient Id.	der	(yrs)) ischemia	Relevant history	Smoking	Locus of occlusion	Age of occlusion	Length of occlusion (cm)
akSTAR(E6 VCL	50,K74 M	O,D82, 69	A,S84A,E108A,F Acute	SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K199A,K130T,K135R,K136A,V137K) (SY118) VCL M 69 Acute Hypertension, ischemic heart disease,	<u>@</u>	Left AF graft	01	4
REN HOL	ΣΣ	69 89	Subacute Acute	Hypercholesterolemia Hypertasion, hypercholesterolemia,	• +	Right PA Right FT bypass	82 92 10 10 10 10 10 10 10 10 10 10 10 10 10	14
PAR MAC	22>	882	Pain, swelling Subacute Acute	Ischemic heart disease, left FP graft Hypertension, ABF graft	• +	Left poplitest to communal fernoral vein Left FP graft Left branch ABF graft	ein 50 30 20	8.4 C
LCT A D/V 2EA	NEW PERE	11 ± 2.7					21 ± 6.9	11 ± 1.8
VERH	7,503,A	52 52	VERH F 52 Claudication	190A,E99D,T101S,E108A,K109A,K130T,K135R,K136A,V137K) (SY141) Hypertension, hypercholesterolemia, right IF endoprothesis	SR,K136, +	A,V137K) (SY141) Right IA	14	12
VAP	ΣΣ	\$ 4	Claudication Claudication	Hypertension, stenting left, right IA Hypertension, hypercholesterolemia,	+ •	Right EIA Aortabifurcation	30 21	18
WYN	Σ	43	Claudication	stenting left + right IA CAD; hypercholesterolemia; stenting left	+	Left FP graft	5.0	30
HOR	ΣΣ	57 25	Acute Acute	FP graft Hypertension; left FP graft Diabetes; hypertension; cardiac valve	+ •	Lefi CIA, left FP graft Left SF artery	7.0	% ≏
Mean ± SEM	EM	55±4.6	9	replacements			.,,,,,	
STAR(K35	4,E650	,K74R	ZA,T	90A,E99D,T101S,E108A,K109A,K130T,K135R,K136A, V137K) (SY145)	R.K136A	V137K) (SY145)	13 I 4.3	19 I 3.3
Z 12	τ Σ	8 6		Hypertension, ischemic heart disease	+	Right SF artery	7.0	91
LAM	Σ		Acide	Hypertension	+	Left PA	21	9.0
BAS	Σ	2	Acute	rr graft heart disease,		Right FP graft Right SFA	7.0	7 2 9 9 9
TOU	-		Acute	hypertension Ischemic heart disease, hypertension	+	Right PA	0.1	0.9
Mean I SEM	ė EM	04 ± 4.	MEAN I SEM 64 ± 4.			ı	36796	12+40

	V147K) or SakS	FAR(K35A,E65Q,	K74R, D82A, S84A,	T90A,E99D,T101S,E108A,K109	KI301, NI33R, NI33R, NI37R, NI37R, D81A, S84A, T90A, E99D, T101S, E108A, KI09A, NI39L, NI33R, NI39R, NI39R, NI37R,
Compound Patient	Recanalization by thrombolysis	Total dose of thrombolytic agent	Total duration of infusion (hrs)	Additional therapy	Complications and remarks
P) din	I A GSELV Tocco	36A (V137K) (SY118)	
SakSTAR(E6	50,K74Q,D82A,S	84A,E108A,K109A,	ALINCLIA, IUCIA,	SakSTAR(E650,K74Q,D82A,S84A,E108A,K109A,N1501,N155M,N1501)	Puncture site hematoma
VCL.	Complete	7	3 5	None	Puncture site hematoms
NEN NEW	Complete	74	4 6	and N	None
HOL	Complete	*	0. 9	•	Small subdural hematoma
PAR	Partial (normal	9	0.0		
	patency with				
	after first control)		;	New FP scaff	None
FRA	Complete		2.7	None	Brain stem hemorrhage; death
MAC	Complete	8.0	0.0		
Mean # SEM		16 ± 2.8	13 # 3.4	V. 100 A V. 120 T K 135R. K 136A.	V137K) (SY141)
SakSTAR(K.	15A,E65Q,K74Q,I	382A,S84A,T90A,E	799D,T1015,E1087	SakSTAR(K35A,E65Q,K74Q,D62A,S84A,T90A,E99D,11015),E108A,N.157A,H.2015,1004	•
VERH	Complete	51	<u>s</u> 5		
DUB	Complete	0.9	J. 6	DIA stenting bilateral IA stenting	Puncture site hematoma
VAP	Complete	4	7 8	EP oraft revision	None
WYN	Complete	ສ	3 3	auo'N	None
HOR	Complete	2	0.5	and N	Retroperitoneal hematoma, died due to septicemia
CNA	Complete	13		2001	
Mean + SEM		14±2.3	16±3.7	A 251 V 1362 V 1364	V137K) (SY145)
SAKSTAR(K	35A,E65Q,K74R,I	D82A,S84A,T90A,E	E99D,T101S,E108,	SakSTAR(K35A,E65Q,K74R,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K109A,K105A,K105A,K105A, SakSTAR(K35A,E65Q,K74R,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K109A,K105A,K10	Retroperitoneal hematoma, Hemorragies
Z	Complete	4	* 3	aco'N	None
DEL	Complete	7.0	0.6	e con	Puncture site hematoma
LAM	Complete	23	2	None	
BAS TOU	Complete				
		15+46	20 ± 7.5		
Mean # SEM		25.4.0			

PTA, percutaneous transluminal angioplasty; IF: iliofemoral; FT: femorotiblal; FP: femoropopliteal.

.12: Neutralizing untibody activity before and after administration of SakSTAR(E65Q,K74Q,D82A,S84A,E108A,K109A,K130T,K135R,K136A, V137K), SakSTAR(K35A,E65Q,K74Q,D82A,S94A,T90A,E99D,T101S,E108A,K136A,K130T,K136R,K136R,K136A,V137K) or SakSTAR(K35A,E65Q,K74R,D82A,S84A,T90A,E99D,T101S,E108A,K130T,K135R,K136A,V137K) in patients with peripheral arterial occlusion	Compound Neutralizing antibody activity (µg/ml) Patient Id.
Table 17:	Comp Pati

	7137K) (SY118)							
4 weeks	9A,K130T,K135R,K136A,	50	0.9	18	15	39	•	18
3 weeks	,S84A,E108A,K10	46	1.6	22	61	15	•	61
Before	650,K740,D82A	0.2	0.1	0.5	0.1	1.2	0.0	0.15
	SakSTAR(E	\CL	REN	HOL	PAR	FRA	MAC	Median

K130T, K135R,K136A,V137K) (SY14							
kSTAR(K35A,E65Q,K74Q,D82A,S84A,T90A,E99D,T101S,E108A,K109A,K1	0.2	2.0	0.0	5.6	0.0	100	1:1
Q,D82A,S84A,T9	0.3	4.3	0.0	01	0.1	1	0.3
35A,E65Q,K74	0.2	0.2	0.0	0.2	0.2	0.8	0.2
SakSTAR(K	VERH	DUB	VAP	WYN	HOR	AND	Median

(<u>)</u>

Table 18: Immunogenicity of SakSTAR variants in patients with peripheral arterial occlusion

	c	Neutralizing activity (us/ml)	activity >5 µg/ml	Specific 1gG (µg/ml)	Code
	8	69 12 (4 - 100)	98	380 (81 - 1850)	
	9	9.0 (0.1 - 23)	3	420 (31 - 730)	SY15
	<u>æ</u>	18 1.5 (0.2 - 7.0)	*	30 (24 - 100)	8Y19
	•	27 (17 - 49)	٠,	2000 (1300 - 3600)	SY118
SakSTAR(E65Q.K/4Q.D82A,S84A,E108A,R109A,R1501,R155R,R156A,C135R,R136A,V137K 6 0.7 (0.1 - 4.3)	9	0.7 (0.1 – 4.3)	7	7.7 (5.1 – 510)	SY141
SakSTAR(K35A,E65Q,K74R,D82A,S84A,190A,E99D,T101S,E108A,K109A,K130T,K135R,K136A,V137K 3 4.7	m	4.7	_		SY145

TROIS IS CYSICING SUBSTITUTION VALIBITIS OF SAKE LAN	VALUATURE OF CA	MALAN					
Variant	Spec. Act. (kU/mg)	Spec. Act. Dimerization level (%) F (kU/mg)	PEG derivatization	Clot lysis in vitro (C _{so} in µg/ml)	t1/2(α) (min)	Clp (ml/min)	Antibody Absorption (Pool 40, %)
SakSTAR	130	0	none	0.33	2.0	2.2	95
SakSTAR (K102C)	143	0	none	0.29	pu	рu	95
SakSTAR (K102C-PEG)	108	0	1	09.0	3.0	0.32	
SakSTAR (K109C) monomeric	001	0	none	0.52	pu	멀	
SakSTAR (K109C) dimeric	1,650	09<	none	0.17	3.6	0.52	06
	2,235	>95	none	0.12	pu	рu	
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Table 20: Cysteine-substitution variants of SakSTAR with reduced immunogenicity, substituted with maleimide-polyethylene glycol

		Specific	Human	Hamsters	රි	Aniibody
Code		activity	plasma	bolus	(ml/mln)	bolus (ml/mln) sbsorption
		(kU/mg)	(kU/mg) (C _w : µg/ml) (C _w : µg/kg)	(C.s.: µg/kg		P40 (%)
		130	0 23	120	2.2	95
•	SetSTAR	3				
		,	3		3.7	2.5
SY19	SakSTAR(E65D.K74R.E80A.D82A.K130T,K135R)	40	Q. K		;	;
	Careta Diese cash Wald Dena Desa King King B	51	0.37	42	0.45	28
31 18(350-373)	URAD 1971 GAST G. FUCULTAT ST. FUCULTATION CO. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	20	0.65		0.28	20
SY19(S3C-MP5)	UBKUTAR(BUCTAR(BUCTAR)	43	0.42	50	0.15	57
SY19(82C-8P5,S3C-SP5)	SEKOLAH(62C;-676,54C;-676,54C;-676,775,775,775,775,775,775,775,775)	90	0.70	8	0.065	57
SY19(S3C-P20)	Sakstar(83C-P20,e65D,ktar,e60A,de2a,ktsot,ktsor)	; ;	9	6	01.0	1.5
SY19(S3C-P10)	SakSTAR(83C-P10,E85D,K74R,E80A,D82A,K130T,K135R)	-	0.00	3	<u>;</u>	
	C-LGTAGNARA EGEN WIND NESA GRAE TONG FOOD THOSE KIDSE KISST KISSE KISSE VISSE (KISSE KISSE)	3,700	0.19		0.95	54
34141 64444637 6861	8A.V137K)	1,200	0.24	12		18
SYTATION SPECIAL SPECI	8A,0137K)	1,400	0.28			18
STIMI(SEC.SPS, SSC-SPS)	DERSTANDING TO BE ARED KTAD FAND TRADE TRADE FROD TIOUS FIDBAK (1994 K130T K135R)	65	0.33	9	0.08	32
ST160(33C-P20)	DORD FAT DO CHARLING CAN THE LOCAL CONTROL OF THE C	7.1	0.36	15	0.56	35
SY161(S3C-MP5)	SOKSTAR(SUG-MPB.KJSA,EDSULN/AR,EGUN,DOKA, PGUN,ESUN, PLOMESTON, PLOMESTON, PLOMESTON, PLOMESTON, PLOMESTON, PLOMESTON PLOMESTO	99	0.40	6	0.15	38
SY181(S3C-P10)	SBKS1 AR(83C-P10, NSSA, E83C, K74R, E80A, D82A, 190A, E89D, 1101S, E108A, K109A, K130T, K135R)	155	0.32	6 0	0.04	4 4

*SP5: OPSS-PEG 5 kDa; MP5: MAL-PEG 5 kDa; P10: MAL-PEG 10 kDa; P20: MAL-PEG 20 kDa.

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CLAIMS

- Staphylokinase derivatives showing a reduced immunogenicity as compared to wild-type staphylokinase,
 after administration to patients with arterial thrombosis.
- Staphylokinase derivatives as claimed in claim 1 having essentially the amino acid sequence as depicted in figure 1 in which one or more amino acids
 have been replaced by another amino acid thus reducing the reactivity with a panel of murine monoclonal antibodies.
- 3. Staphylokinase derivatives as claimed in claim 1 having essentially the amino acid sequence as depicted in figure 1 in which one or more amino acids have been replaced by another amino acid thus reducing the absorption of SakSTAR-specific antibodies from plasma of patients treated with staphylokinase.
- 4. Staphylokinase derivatives as claimed in
 20 claim 1 having essentially the amino acid sequence as
 depicted in figure 1 in which one or more amino acids
 have been replaced by other amino acids, without reducing
 the specific activity by more than 50 percent.
- 5. Staphylokinase derivatives SakSTAR(K35X, G36X,E65X,K74X,E80X,D82X,K102X,E108X,K109X,K121X,K130X,K135X,K136X,+137X) having the amino acid sequence as depicted in figure 1 in which one or more of the amino acids Lys in position 35, Gly in position 36, Glu in position 65, Lys in position 74, Glu in position 80, Asp in position 82, Lys in position 102, Glu in position 108, Lys in position 109, Lys in position 121, Lys in position 130, Lys in position 135 and/or Lys in position 136 have been replaced with other amino acids and/or in which one amino acid has been added at the COOH-terminus, thus altering the immunogenicity after administration in patients, without markedly reducing the specific activity.

7. Staphylokinase derivative as claimed in

- 6. Staphylokinase derivatives listed in Tables 1, 3, 4, 5, 6, 7, 8, 13, 19 and 20, having the amino acid sequence as depicted in figure 1 in which the indicated amino acids have been replaced by other amino acids thus reducing the absorption of SakSTAR-specific antibodies from plasma of patients treated with staphylokinase, without reducing the specific activity.
- claims 1-6 selected from the group consisting of

 10 SakSTAR(K74A,E75A,R77A), SakSTAR(K35A,E75A),

 SakSTAR(E75A), SakSTAR(E80A,D82A), SakSTAR(E80A),

 SakSTAR(D82A), SakSTAR(E75A,D82A), SakSTAR(S34G,G36R,

 H43R), SakSTAR(K35A), SakSTAR(D82A), SakSTAR(D82A,S84A),

 SakSTAR(T90A), SakSTAR(Y92A), SakSTAR(K130A),
- 15 SakSTAR(V132A), SakSTAR(S34G,G36R,H43R), SakSTAR(G36R), SakSTAR(H43R), SakSTAR(G36R,K74R), SakSTAR(K35E), SakSTAR(K74Q), SakSTAR(K130T), SakSTAR(V132L), SakSTAR(V132T), SakSTAR(V132N), SakSTAR(V132R), SakSTAR(K130T,K135R),
- 20 SakSTAR(K74R,K130T,K135R), SakSTAR(K74Q,K130T,K135R), SakSTAR(G36R,K74R,K130T,K135R), SakSTAR(G36R,K74Q,K130T,K135R), SakSTAR(G36R,H43R,K74R,K130T,K135R), SakSTAR(E65A,K74Q,K130T,K135R), SakSTAR(E65Q,K74Q,K130T,K135R), SakSTAR(K74Q,K86A,K130T,K135R),
- 25 SakSTAR(E65Q,T71S,K74Q,K130T,K135R), SakSTAR(K74Q, K130A,K135R), SakSTAR(E65Q,K74Q,K130A,K135R), SakSTAR(K74Q,K130E,K135R), SakSTAR(K74Q,K130E, V132R,K135R), SakSTAR(E65Q,K74Q,T90A,K130A,K135R), SakSTAR(E65Q,K74Q,N95A,K130A,K135R), SakSTAR(E65Q,K74Q,
- 30 E118A,K130A,K135R), SakSTAR(E65Q,K74Q,N95A,E118A,K130A,K135R), SakSTAR(N95A,K130A,K135R), SakSTAR(E65Q,K74Q,K109A,K130,K135R), SakSTAR(E65Q,K74Q,E108A,K109A,K130T,K135R), SakSTAR(E65Q,K74Q,K121A,K130T,K135R), SakSTAR(E65Q,K74Q,K121A,K130T,K135R),
- 35 SakSTAR(E80A, D82A, K130T, K135R), SakSTAR(K74R, E80A, D82A, K130T, K135R), SakSTAR(K74Q, E80A, D82A, K130T, K135R), SakSTAR(K35A, K74R, E80A, D82A, K130T, K135R), SakSTAR(E65D, K74R, E80A, D82A, K130T, K135R), SakSTAR(E65S, K74R, E80A,

D82A,K130T,K135R), SakSTAR(S34G,G36R,K74R,K130T,K135R), SakSTAR(E65A,K74R,E80A,D82A,K130T,K135R), SakSTAR(E65N, K74R,E80A,D82A,K130T,K135R), SakSTAR(E65Q,K74R,E80A, D82A,K130T,K135R), SakSTAR(K57A,E58A,E61A,E80A,D82A,

- 5 K130T,K135R), SakSTAR(E65D,K74Q,E80A,D82A,K130T,K135R), SakSTAR(E65Q,K74Q,E80A,D82A,K130T,K135R), SakSTAR(K35A, E65D,K74Q,E80A,D82A,K130T,K135R), SakSTAR(K74R,E80A,D82A, S103A,K130T,K135R), SakSTAR(E65D,K74R,E80A,D82A,K109A, K130T,K135R), SakSTAR(E65D,K74R,E80A,D82A,K130T,
- 10 K135R,K136A), SakSTAR(E65Q,K74Q,D82A,S84A,K130T,K135R),
 SakSTAR(K35A,K74Q,E80A,D82A,K130T,K135R), SakSTAR(K35A,
 E65D,K74R,E80A,D82A,K130T,K135R).
 - 8. SakSTAR(E65D, K74R, E80A, D82A, K130T, K135R) having the code SY19.
- 9. SakSTAR(K35A,E65Q,K74R,E80A,D82A,T90A,E99D, T101S,E108A,K109A,K130T,K135R) having the code SY161.
 - 10. Staphylokinase derivatives as claimed in claims 1-9 having an amino acid substituted with Cys, resulting in dimerization and/or increased specific
- 20 activity and/or reduced clearance and/or increased thrombolytic potency.
- 11. Staphylokinase derivatives as claimed in claims 1-10 with polyethylene glycol substitution, characterized by a maintained specific activity and a significantly reduced plasma clearance.
 - 12. Staphylokinase derivatives as claimed in claim 10 wherein the Cys is chemically modified with polyethylene glycol with molecular weights up to 20 kDa.
- 13. Staphylokinase derivatives as claimed in claim 12 wherein selected amino acids in the NH₂-terminal region of 10 amino acids, are substituted with Cys, which is chemically modified with polyethylene glycol with molecular weights up to 20 kDa, which derivatives are characterized by a significantly reduced plasma clearance
- 35 and maintained thrombolytic potency upon single intravenous bolus administration at a reduced dose.
 - 14. Staphylokinase derivative as claimed in claim 13, wherein the serine in position 2 or 3 is

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substituted with a cystein and the cystein is chemically modified with polyethylene glycol having a molecular weight of 5, 10 or 20 kDa.

- 15. Staphylokinase derivative as claimed in 5 claim 14, which derivative is SY161(S3C-MP5) as defined in table 20.
 - 16. Staphylokinase derivative as claimed in claim 14, which derivative is SY161(S3C-P10) as defined in table 20.
- 17. Staphylokinase derivative as claimed in claim 14, which derivative is SY161(S3C-P20) as defined in table 20.
- 18. Staphylokinase derivative as claimed in claim 14, which derivative is SY19(S3C-MP5) as defined in 15 table 20.
 - 19. Staphylokinase derivative as claimed in claim 14, which derivative is SY19(S3C-SP5) as defined in table 20.
- 20. Staphylokinase derivative as claimed in 20 claim 14, which derivative is SY19(S2C-SP5,S3C-SP5) as defined in table 20.
 - 21. Staphylokinase derivative as claimed in claim 14, which derivative is SY19(S3C-P20) as defined in table 20.
- 25 22. Staphylokinase derivative as claimed in claim 14, which derivative is SY19(S3C-P10) as defined in table 20.
 - 23. Dimer of two staphylokinase derivatives as claimed in claim 10.
- 24. Method for producing the staphylokinase derivatives as claimed in claims 1 to 10, comprising the steps of:
- a. preparing a DNA fragment comprising at least the part of the coding sequence of staphylokinase that
 35 provides for its biological activity;
 - b. performing in vitro site-directed mutagenesis on the DNA fragment to replace one or more

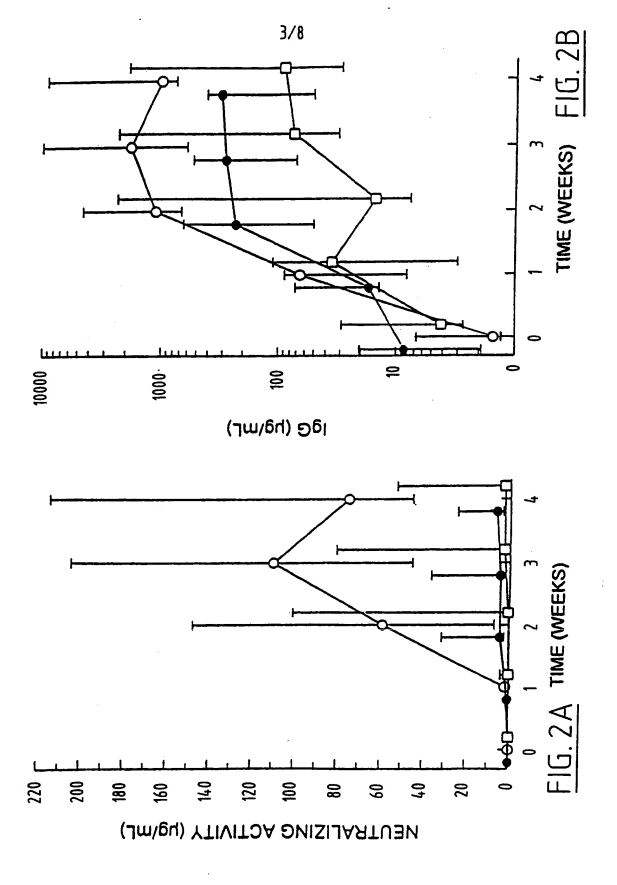
codons for wild-type amino acids by a codon for another amino acid;

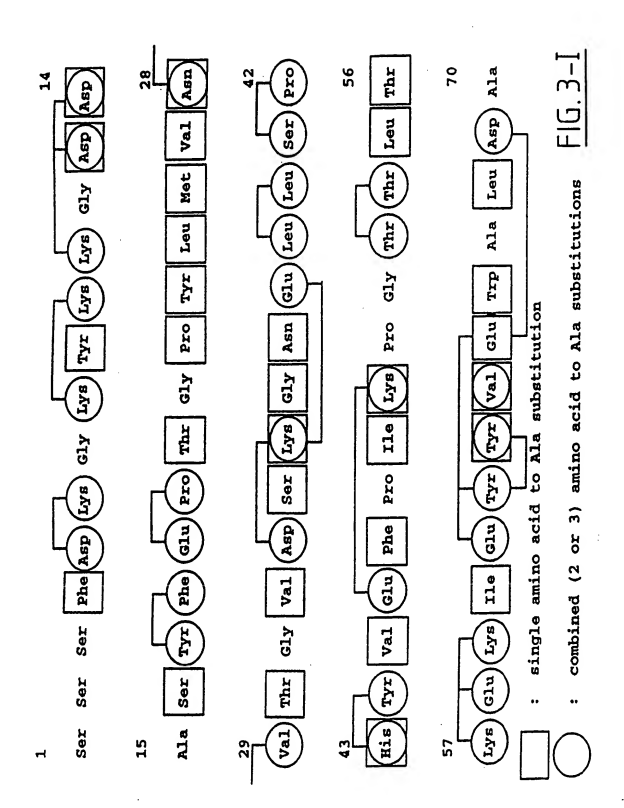
- c. cloning the mutated DNA fragment in a suitable vector;
- d. transforming or transfecting a suitable host cell with the vector; and
 - e. culturing the host cell under conditions suitable for expressing the DNA fragment.
- 25. Method as claimed in claim 24, wherein the 10 DNA fragment is a 453 bp EcoRI-HindIII fragment of the plasmid pMEX602sakB, the <u>in vitro</u> site-directed mutagenesis is performed and the mutated DNA fragment is expressed in <u>E. coli</u>.
- 26. Pharmaceutical composition comprising at
 15 least one of the staphylokinase derivatives as claimed in
 claims 1 to 23 together with a suitable excipient.
 - 27. Pharmaceutical composition as claimed in claim 26 for treating arterial thrombosis.

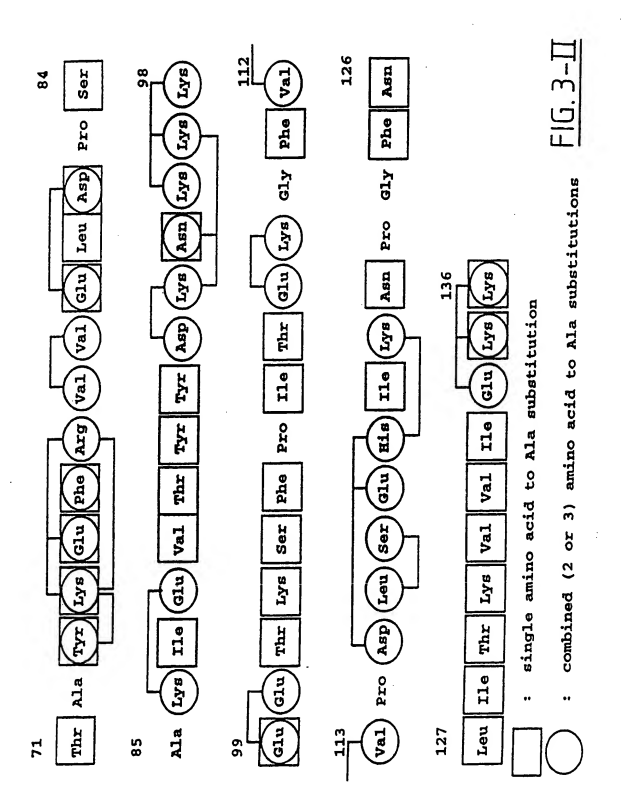
				1/8	}				
14 Asp	78	Asn	42	Pro	26	Thr	70	Ala	
AGD A		Val		Ser		Leu		Asp	
Gly A		Met		Leu		Thr		Leu	
Lys G		ren		Leu		Thr		Ala	
Lys I	·	Tyr		Glu		Gly		Trp	
Tyr 1		Pro		Asn		Pro		Glu	
Lys '		Gly		Gly		Lys		Val	
Gly		Thr		Lys		116		Tyr	FIG. 1-I
Lys		Pro		Ser		Pro		Tyr	
ABP		Glu		Asp		Phe		Glu	
Phe		Phe		Val		Glu		110	
Ser		TYT		Gly		Val		Lys	
Ser		Ser		Thr		Tyr		Glu	

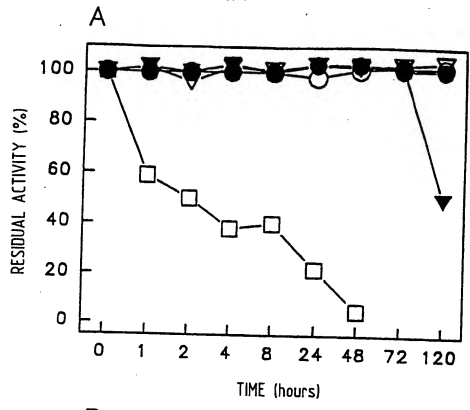
				2/	8				
84 Ser	86	Lys	112	Val	126	Asn			
Pro		Lys		Phe		Phe			
Asp		Lys		$\mathbf{G}1\mathbf{y}$		Gly			
Leu		Asn		Lys		Pro			
Glu		Lys		Glu		Asn	136	Lys	
Val		Asp		Thr		Lys		Lys	
Val		Tyr		Ile		110		Ile Glu	믜
Arg		TY		Pro		His			FIG. 1-I
Phe		Thr		Phe		Glu		Val	
Glu		Val		Ser		Ser	•	Val	
Lys		Glu		Lys		Leu		Lys	
Tyr		116		Thr		Asp		Thr	
Ala		Lys		Glu		Pro		Ile	
71 rhr	82	A.La	66	Glu	113	Val	127	Leu	

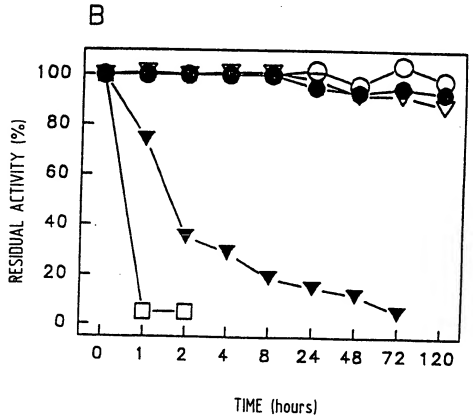
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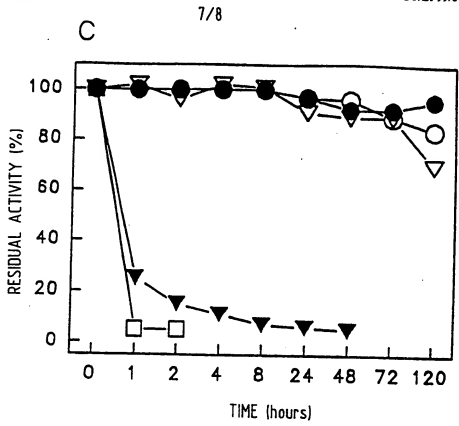


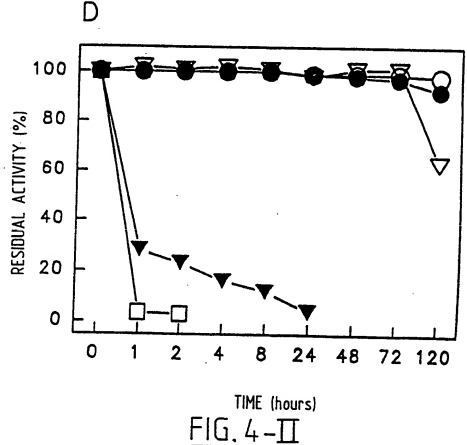


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FIG. 4-I

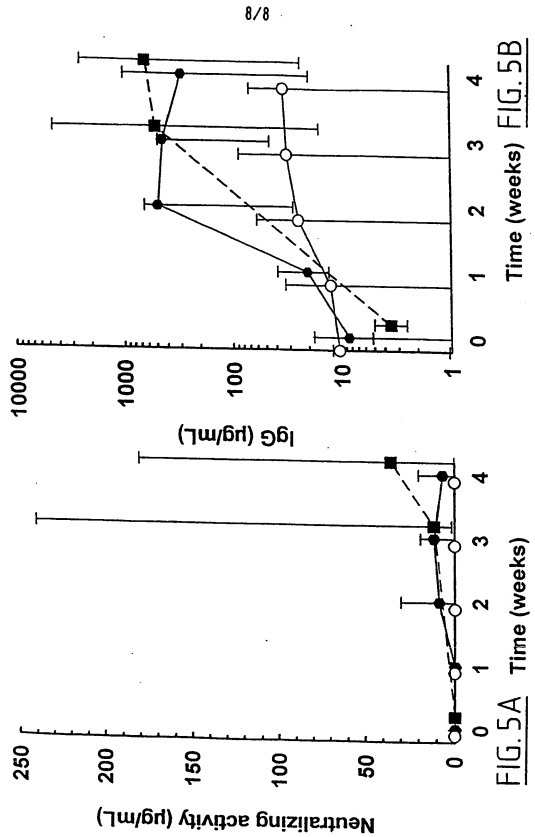
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